Food Lipids
Chemistry, Nutrition, and Biotechnology
Second Edition, Revised and Expanded

edited by
Casimir C. Akoh
The University of Georgia
Athens, Georgia

David B. Min
The Ohio State University
Columbus, Ohio

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Preface to the Second Edition

Readers’ responses to the first edition, published in 1998, were overwhelming, and we are grateful. The response reassured us that there was indeed great need for a textbook suitable for teaching food lipids, nutritional aspects of lipids, and lipid chemistry courses to food science and nutrition majors. The aim of the first edition remains unchanged: to provide a modern, easy-to-read textbook for students and instructors. The book is also suitable for upper-level undergraduate, graduate, and postgraduate instruction. Scientists who have left the university and are engaged in research and development in the industry, government, or academics will find this book a useful reference. Again, we made every effort to select contributors who are internationally recognized experts. We thank them for their exceptional attention to details and timely submissions.

The text has been updated with new information. The indexing has been improved. We changed the order of chapters and added two new chapters, “Conjugated Linoleic Acid” and “Food Applications of Lipids.” While it is not possible to cover every aspect of lipids, we feel we have added and covered most topics that are of interest to our readers. The book still is divided into five main parts: Chemistry and Properties; Processing; Oxidation; Nutrition; and Biotechnology and Biochemistry.

Obviously, we made some mistakes in the first edition. Thanks go to our students for pointing out most of the obvious and glaring errors. Based on readers’ and reviewers’ comments, we have improved the new edition—we hope without creating new errors, which are sometimes unavoidable for a book this size and complexity. We apologize for any errors and urge you to contact us if you find mistakes or have suggestions to improve the readability and comprehension of this text.
Special thanks to our readers and students, and to the editorial staff of Marcel Dekker, Inc., for their helpful suggestions toward improving the quality of this edition.

Casimir C. Akoh
David B. Min
Preface to the First Edition

There is a general consensus on the need for a comprehensive, modern textbook of food lipids that will provide a guide to students and instructors, as well as cover the topics expected of students taking a food lipids or lipid chemistry course as food science and nutrition majors. The text is suitable for undergraduate and graduate instruction. In addition, food industry professionals seeking background or advanced knowledge in lipids will find this book helpful. It is envisaged that this book will also serve as a reference source for individuals engaged in food research, product development, food processing, nutrition and dietetics, quality assurance, oil processing, fat substitutes, genetic engineering of oil crops, and lipid biotechnology. It is expected that students and others using this book will have backgrounds in chemistry and biochemistry.

Every effort was made to involve internationally recognized experts as contributors to this text. Considerable efforts were made by the authors to start from basics and build up and to provide copious equations, tables, illustrations, and figures to enhance teaching, comprehension, and to drive the lecture materials home. Mechanisms of reactions are given to help in the understanding of the underlying principles of lipid chemistry and hopefully will lead to solutions of adverse reactions of lipids in the future. We believe that the end product of this work provides state-of-the-art and authoritative information that covers almost all aspects of food lipids and will serve as a unique text for instruction throughout the world. The text is reader-friendly and easy to understand. Adequate references are provided to encourage persons who need to inquire further or need detailed information on any aspect covered in this book.
The text is divided into five main parts, namely: Chemistry and Properties; Processing; Oxidation; Nutrition; and Biotechnology and Biochemistry.

Part I is devoted to introductory chapters on the nomenclature and classification of lipids, chemistry of phospholipids, waxes and sterols, emulsion and emulsifiers, frying, and on the analysis of lipids including \textit{trans} fatty acids. It is important to understand the structure and chemistry of lipids and some basic concepts before moving on to more complex and applied topics.

Part II deals with the technology of edible oils and fats processing including refining, recovery, crystallization, polymorphism, chemical interesterification, and hydrogenation.

Part III describes the key oxidation reactions in both edible oils and plant and animal or muscle tissues. Lipid oxidation is a major cause of quality deterioration of processed and unprocessed foods. Methods to measure lipid oxidation in fats and oils are given. The mechanism of antioxidant actions in arresting or improving the oxidative stability of foods is discussed. This section has tremendous implications for food technologists and nutritionists as the consumer continues to demand and expect nothing but high-quality foods and food products.

Part IV deals with the role of fats and oils in overall nutrition. The importance of antioxidants in nutrition and food preservation is presented. Excess fat intake is associated with many disease conditions. This section describes various omega fatty acids and their sources, the role of dietary fats in atherosclerosis, eicosanoids production, immune system, coronary heart disease and obesity. The various types of lipid-based synthetic fat substitutes are discussed.

Part V introduces the new biotechnology as applied to lipids and production of value-added lipid products. The microbial lipases used in enzyme biotechnology are discussed. The potential for replacing chemical catalysis with enzyme catalysis are described further in the chapters dealing with enzymatic interesterification and structured lipids. Lipid biotechnology and biosynthesis chapters set the stage for a better understanding of the chapter on genetic engineering of plants that produce vegetable oil and for further research in lipid biotechnology, a dynamic area of increasing industrial interest.

We feel that we covered most of the topics expected for a food lipids course in this text. It is hoped that this edition will stimulate discussions and generate helpful comments to improve upon future editions. Unavoidably, in a book of this size and complexity, there are some areas of overlap. Efforts are made to cross reference the chapters as such.

Finally, we would like to thank all the authors for the magnificent work they did in making sure that their contributions are timely, easy to read, and most of all, for their time and devotion to details and accuracy of information presented. The help of the Marcel Dekker editorial staff is greatly appreciated, with special thanks to Rod Learmonth and Maria Allegra.

\textit{Casimir C. Akoh}

\textit{David B. Min}
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Contributors

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    Kirk L. Parkin

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    Vic C. Knauf and Anthony J. Del Vecchio
Contributors

Casimir C. Akoh  Department of Food Science and Technology, The University of Georgia, Athens, Georgia

M. Margaret Barth  Redi-Cut Foods, Inc., Franklin Park, Illinois

Jeffrey M. Boff  Department of Food Science and Technology, The Ohio State University, Columbus, Ohio

Terrence L. Boos  Department of Chemistry, Auburn University, Auburn, Alabama

Eric A. Decker  Department of Food Science, University of Massachusetts, Amherst, Massachusetts

Anthony J. Del Vecchio  Monsanto Inc., Davis, California

Paul S. Dimick  Department of Food Science, The Pennsylvania State University, University Park, Pennsylvania

Ronald R. Eitenmiller  Department of Food Science and Technology, The University of Georgia, Athens, Georgia

Marilyn C. Erickson  Center for Food Safety, Department of Food Science and Technology, The University of Georgia, Griffin, Georgia
J. Bruce German  Department of Food Science and Technology, University of California, Davis, California

Barbara Mullen Grossman  Department of Foods and Nutrition, The University of Georgia, Athens, Georgia

Frank D. Gunstone  Scottish Crop Research Institute, Dundee, Scotland

Dorothy B. Hausman  Department of Foods and Nutrition, The University of Georgia, Athens, Georgia

Dana R. Higbee  Department of Foods and Nutrition, The University of Georgia, Athens, Georgia

David Hildebrand  Department of Agronomy, University of Kentucky, Lexington, Kentucky

Lawrence A. Johnson  Center for Crops Utilization Research, Department of Food Science and Human Nutrition, Iowa State University, Ames, Iowa

David M. Klurfeld  Department of Nutrition and Food Science, Wayne State University, Detroit, Michigan

Vic C. Knauf  Monsanto Inc., Davis, California

David Kritchevsky  The Wistar Institute, Philadelphia, Pennsylvania

Patrick J. Lawler  McCormick and Company Inc., Hunt Valley, Maryland

Shengrong Li  Department of Chemistry, Auburn University, Auburn, Alabama

Yong Li  Center for Enhancing Foods to Protect Health, Purdue University, West Lafayette, Indiana

Dorris A. Lillard  Department of Food Science and Technology, The University of Georgia, Athens, Georgia

Alejandro G. Marangoni  Department of Food Science, University of Guelph, Guelph, Ontario, Canada

D. Julian McClements  Department of Food Science, University of Massachusetts, Amherst, Massachusetts

Richard E. McDonald  Division of Food Processing and Packaging, National Center for Food Safety and Technology, U.S. Food and Drug Administration, Summit-Argo, Illinois

Ronald P. Mensink  Department of Human Biology, Maastricht University, Maastricht, The Netherlands
David B. Min  Department of Food Science and Technology, The Ohio State University, Columbus, Ohio

Magdi M. Mossoba  Center for Food Safety and Applied Nutrition, U.S. Food and Drug Administration, Washington, D.C.

Kumar D. Mukherjee  Institute for Biochemistry and Technology of Lipids, H. P. Kaufmann-Institute, Federal Center for Cereal, Potato, and Lipid Research, Munster, Germany

Sean Francis O’Keefe  Department of Food Science and Technology, Virginia Polytechnic Institute and State University, Blacksburg, Virginia

Edward J. Parish  Department of Chemistry, Auburn University, Auburn, Alabama

Kirk L. Parkin  Department of Food Science, University of Wisconsin–Madison, Madison, Wisconsin

Jogchum Plat  Department of Human Biology, Maastricht University, Maastricht, The Netherlands

David W. Reische  The Dannon Company, Inc., Fort Worth, Texas

Dérick Rousseau  School of Nutrition, Ryerson University, Toronto, Ontario, Canada

Fereidoon Shahidi  Department of Biochemistry, Memorial University of Newfoundland, St. John’s, Newfoundland, Canada

Elisabeth H. M. Temme  Department of Public Health, University of Leuven, Leuven, Belgium

P. K. J. P. D. Wanasundara  Department of Applied Microbiology and Food Science, University of Saskatchewan, Saskatoon, Saskatchewan, Canada

Udaya N. Wanasundara  Pilot Plant Corporation, Saskatoon, Saskatchewan, Canada


Bruce A. Watkins  Center for Enhancing Foods to Protect Health, Purdue University, West Lafayette, Indiana

Steven M. Watkins  Lipomics Technologies, Inc., West Sacramento, California

John D. Weete  West Virginia University, Morgantown, West Virginia

Wendy M. Willis  Ives’ Veggie Cuisine, Vancouver, British Columbia, Canada

Hong Zhuang  Redi-Cut Foods, Inc., Franklin Park, Illinois
I. DEFINITIONS OF LIPIDS

No exact definition of lipids exists. Christie [1] defines lipids as “a wide variety of natural products including fatty acids and their derivatives, steroids, terpenes, carotenoids and bile acids, which have in common a ready solubility in organic solvents such as diethyl ether, hexane, benzene, chloroform or methanol.”

Kates [2] says that lipids are “those substances which are (a) insoluble in water; (b) soluble in organic solvents such as chloroform, ether or benzene; (c) contain long-chain hydrocarbon groups in their molecules; and (d) are present in or derived from living organisms.”

Gurr and James [3] point out that the standard definition includes “a chemically heterogeneous group of substances, having in common the property of insolubility in water, but solubility in non-polar solvents such as chloroform, hydrocarbons or alcohols.”

Despite common usage, definitions based on solubility have obvious problems. Some compounds that are considered lipids, such as C1–C4 very short chain fatty acids (VSCFAs), are completely miscible with water and insoluble in nonpolar solvents. Some researchers have accepted this solubility definition strictly and exclude C1–C3 fatty acids in a definition of lipids, keeping C4 (butyric acid) only because of its presence in dairy fats. Additionally, some compounds that are considered lipids, such as some trans fatty acids (those not derived from bacterial hydrogenation), are not derived directly from living organisms. The development of synthetic acaloric and reduced calorie lipids complicates the issue because they may fit into solubility-
based definitions but are not derived from living organisms, may be acaloric, and may contain esters of VSCFAs.

The traditional definition of total fat of foods used by the U.S. Food and Drug Administration (FDA) has been “the sum of the components with lipid characteristics that are extracted by Association of Official Analytical Chemists (AOAC) methods or by reliable and appropriate procedures.” The FDA has changed from a solubility-based definition to “total lipid fatty acids expressed as triglycerides” [4], with the intent to measure caloric fatty acids. Solubility and size of fatty acids affect their caloric values. This is important for products that take advantage of this, such as Benefat/Salatrim, so these products would be examined on a case-by-case basis. Food products containing sucrose polyesters would require special methodology to calculate caloric fatty acids. Foods containing vinegar (~4.5% acetic acid) present a problem because they will be considered to have 4.5% fat unless the definition is modified to exclude water-soluble fatty acids or the caloric weighting for acetic acid is lowered [4].

Despite the problems with accepted definitions, a more precise working definition is difficult given the complexity and heterogeneity of lipids. This chapter introduces the main lipid structures and their nomenclature.

II. LIPID CLASSIFICATIONS

Classification of lipid structures is possible based on physical properties at room temperature (oils are liquid and fats are solid), their polarity (polar and neutral lipids), their essentiality for humans (essential and nonessential fatty acids), or their structure (simple or complex). Neutral lipids include fatty acids, alcohols, glycerides, and sterols, while polar lipids include glycerophospholipids and glyceroglycolipids. The separation into polarity classes is rather arbitrary, as some short chain fatty acids are very polar. A classification based on structure is, therefore, preferable.

Based on structure, lipids can be classified as derived, simple, or complex. The derived lipids include fatty acids and alcohols, which are the building blocks for the simple and complex lipids. Simple lipids, composed of fatty acids and alcohol components, include acylglycerols, ether acylglycerols, sterols, and their esters and wax esters. In general terms, simple lipids can be hydrolyzed to two different components, usually an alcohol and an acid. Complex lipids include glycerophospholipids (phospholipids), glyceroglycolipids (glycolipids), and sphingolipids. These structures yield three or more different compounds on hydrolysis.

The fatty acids constitute the obvious starting point in lipid structures. However, a short review of standard nomenclature is appropriate. Over the years, a large number of different nomenclature systems have been proposed [5]. The resulting confusion has led to a need for nomenclature standardization. The International Union of Pure and Applied Chemists (IUPAC) and International Union of Biochemistry (IUB) collaborative efforts have resulted in comprehensive nomenclature standards [6], and the nomenclature for lipids has been reported [7–9]. Only the main aspects of the standardized IUPAC nomenclature relating to lipid structures will be presented; greater detail is available elsewhere [7–9].

Standard rules for nomenclature must take into consideration the difficulty in maintaining strict adherence to structure-based nomenclature and elimination of common terminology [5]. For example, the compound known as vitamin K₁ can be
described as 2-methyl-3-phytyl-1,4-naphthoquinone. Vitamin \( K_1 \) and many other trivial names have been included into standardized nomenclature to avoid confusion arising from long chemical names. Standard nomenclature rules will be discussed in separate sections relating to various lipid compounds.

Fatty acid terminology is complicated by the existence of several different nomenclature systems. The IUPAC nomenclature, common (trivial) names, and shorthand (\( \omega \)) terminology will be discussed. As a lipid class, the fatty acids are often called free fatty acids (FFAs) or nonesterified fatty acids (NEFAs). IUPAC has recommended that fatty acids as a class be called fatty acids and the terms FFA and NEFA eliminated [6].

### A. Standard IUPAC Nomenclature of Fatty Acids

In standard IUPAC terminology [6], the fatty acid is named after the parent hydrocarbon. Table 1 lists common hydrocarbon names. For example, an 18-carbon carboxylic acid is called octadecanoic acid, from octadecane, the 18-carbon aliphatic hydrocarbon. The name octadecanecarboxylic acid may also be used, but it is more cumbersome and less common. Table 2 summarizes the rules for hydrocarbon nomenclature.

Double bonds are designated using the \( \Delta \) configuration, which represents the distance from the carboxyl carbon, naming the carboxyl carbon number 1. A double bond between the 9th and 10th carbons from the carboxylic acid group is a \( \Delta^9 \) bond. The hydrocarbon name is changed to indicate the presence of the double bond. An 18-carbon fatty acid with one double bond to an octadecenoic acid, one with two double bonds octadecadienoic acid, and so on. The double-bond positions are des-

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</tr>
<tr>
<td>16</td>
<td>Hexadecane</td>
<td>70</td>
<td>Heptacosane</td>
</tr>
<tr>
<td>17</td>
<td>Heptadecane</td>
<td>80</td>
<td>Octacosane</td>
</tr>
<tr>
<td>18</td>
<td>Octadecane</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 2  IUPAC Rules for Hydrocarbon Nomenclature

1. Saturated unbranched acyclic hydrocarbons are named with a numerical prefix and the termination “ane.” The first four in this series use trivial prefix names (methane, ethane, propane, and butane), whereas the rest use prefixes that represent the number of carbon atoms.

2. Saturated branched acyclic hydrocarbons are named by prefixing the side chain designation to the name of the longest chain present in the structure.

3. The longest chain is numbered to give the lowest number possible to the side chains, irrespective of the substituents.

4. If more than two side chains are present, they can be cited either in alphabetical order or in order of increasing complexity.

5. If two or more side chains are present in equivalent positions, the one assigned the lowest number is cited first in the name. Order can be based on alphabetical order or complexity.

6. Unsaturated unbranched acyclic hydrocarbons with one double bond have the “ane” replaced with “ene.” If there is more than one double bond, the “ane” is replaced with “diene,” “triene,” “tetraene,” etc. The chain is numbered to give the lowest possible number to the double bond(s).

Source: Ref. 6.

Ignored with numbers before the fatty acid name (Δ9-octadecenoic acid or simply 9-octadecenoic acid). The Δ is assumed and often not placed explicitly in structures.

Double-bond geometry is designated with the cis–trans or E/Z nomenclature systems [6]. The cis/trans terms are used to describe the positions of atoms or groups connected to doubly bonded atoms. They can also be used to indicate relative positions in ring structures. Atoms/groups are cis or trans if they lie on the same (cis) or opposite (trans) sides of a reference plane in the molecule. Some examples are shown in Figure 1. The prefixes cis and trans can be abbreviated as c and t in structural formulas.

The cis/trans configuration rules are not applicable to double bonds that are terminal in a structure or to double bonds that join rings to chains. For these conditions, a sequence preference ordering must be conducted. Since cis/trans nomenclature...
Table 3  A Summary of Sequence Priority Rules for E/Z Nomenclature

1. Higher atomic number precedes lower.
2. For isotopes, higher atomic mass precedes lower.
3. If the atoms attached to one of the double-bonded carbons are the same, proceed outward concurrently until a point of difference is reached considering atomic mass and atomic number.
4. Double bonds are treated as if each bonded atom is duplicated. For example:

\[
\begin{align*}
-\text{HC}==\text{CH} & = \quad | \quad | \\
\text{C} & \quad \text{C} \\
-\text{HC}==\text{O} & = \quad | \quad | \\
\text{O} & \quad \text{C}
\end{align*}
\]

Source: Ref. 10.

clature is applicable only in some cases, a new nomenclature system was introduced by the Chemical Abstracts Service and subsequently adopted by IUPAC (the E/Z nomenclature). This system was developed as a more applicable system to describe isomers by using sequence ordering rules, as is done using the R/S system (rules to decide which ligand has priority). The sequence-rule-preferred atom/group attached to one of a pair of doubly bonded carbon atoms is compared to the sequence-rule-preferred atom/group of the other of the doubly bonded carbon atoms. If the preferred atom/groups are on the same side of the reference plane, it is the Z configuration. If they are on the opposite sides of the plane, it is the E configuration. Table 3 summarizes some of the rules for sequence preference [10]. Although \textit{cis} and Z (or \textit{trans} and E) do not always refer to the same configurations, for most fatty acids \textit{E} and \textit{trans} are equivalent, as are \textit{Z} and \textit{cis}.

B. Common (Trivial) Nomenclature of Fatty Acids

Common names have been introduced throughout the years and, for certain fatty acids, are a great deal more common than standard (IUPAC) terminology. For example, oleic acid is much more common than \textit{cis}-9-octadecenoic acid. Common names for saturated and unsaturated fatty acids are illustrated in Tables 4 and 5. Many of the common names originate from the first identified botanical or zoological origins for those fatty acids. Myristic acid is found in seed oils from the Myristicaceae family. Mistakes have been memorialized into fatty acid common names; margaric acid (heptadecanoic acid) was once incorrectly thought to be present in margarine. Some of the common names can pose memorization difficulties, such as the following combinations: caproic, caprylic, and capric; arachidic and arachidonic; linoleic, linolenic, \(\gamma\)-linolenic, and dihomo-\(\gamma\)-linolenic. Even more complicated is the naming of EPA, or eicosapentaenoic acid, usually meant to refer to \(c-5,\ c-8,\ c-11,\ c-14,\ c-17\)-eicosapentaenoic acid, a fatty acid found in fish oils. However, a different isomer \(c-2,\ c-5,\ c-8,\ c-11,\ c-14\)-eicosapentaenoic acid is also found in nature. Both can be referred to as “eicosapentaenoic” acids using standard nomenclature. Nevertheless, in common nomenclature EPA refers to the \(c-5,\ c-8,\ c-11,\ c-14,\ c-17\) iso-
Table 4  Systematic, Common, and Shorthand Names of Saturated Fatty Acids

<table>
<thead>
<tr>
<th>Systematic name</th>
<th>Common name</th>
<th>Shorthand</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methanoic</td>
<td>Formic</td>
<td>1:0</td>
</tr>
<tr>
<td>Ethanoic</td>
<td>Acetic</td>
<td>2:0</td>
</tr>
<tr>
<td>Propanoic</td>
<td>Propionic</td>
<td>3:0</td>
</tr>
<tr>
<td>Butanoic</td>
<td>Butyric</td>
<td>4:0</td>
</tr>
<tr>
<td>Pentanoic</td>
<td>Valeric</td>
<td>5:0</td>
</tr>
<tr>
<td>Hexanoic</td>
<td>Caproic</td>
<td>6:0</td>
</tr>
<tr>
<td>Heptanoic</td>
<td>Enanthic</td>
<td>7:0</td>
</tr>
<tr>
<td>Octanoic</td>
<td>Caprylic</td>
<td>8:0</td>
</tr>
<tr>
<td>Nonanoic</td>
<td>Pelargonic</td>
<td>9:0</td>
</tr>
<tr>
<td>Decanoic</td>
<td>Capric</td>
<td>10:0</td>
</tr>
<tr>
<td>Undecanoic</td>
<td>—</td>
<td>11:0</td>
</tr>
<tr>
<td>Dodecanoic</td>
<td>Lauric</td>
<td>12:0</td>
</tr>
<tr>
<td>Tridecanoic</td>
<td>—</td>
<td>13:0</td>
</tr>
<tr>
<td>Tetradecanoic</td>
<td>Myristic</td>
<td>14:0</td>
</tr>
<tr>
<td>Pentadecanoic</td>
<td>—</td>
<td>15:0</td>
</tr>
<tr>
<td>Hexadecanoic</td>
<td>Palmitic</td>
<td>16:0</td>
</tr>
<tr>
<td>Heptadecanoic</td>
<td>Margaric</td>
<td>17:0</td>
</tr>
<tr>
<td>Octadecanoic</td>
<td>Stearic</td>
<td>18:0</td>
</tr>
<tr>
<td>Nonadecanoic</td>
<td>—</td>
<td>19:0</td>
</tr>
<tr>
<td>Eicosanoic</td>
<td>Arachidic</td>
<td>20:0</td>
</tr>
<tr>
<td>Docosanoic</td>
<td>Behenic</td>
<td>22:0</td>
</tr>
<tr>
<td>Tetraicosanoic</td>
<td>Lignoceric</td>
<td>24:0</td>
</tr>
<tr>
<td>Hexacosanoic</td>
<td>Cerotic</td>
<td>26:0</td>
</tr>
<tr>
<td>Octacosanoic</td>
<td>Montanic</td>
<td>28:0</td>
</tr>
<tr>
<td>Tricontanoic</td>
<td>Melissic</td>
<td>30:0</td>
</tr>
<tr>
<td>Dotriacontanoic</td>
<td>Lacceroic</td>
<td>32:0</td>
</tr>
</tbody>
</table>

mer. Docosahexaenoic acid (DHA) refers to all-cis 4,7,10,13,16,19-docosahexaenoic acid.

C. Shorthand (ω) Nomenclature of Fatty Acids

Shorthand (ω) identifications of fatty acids are found in common usage. The shorthand designation is the carbon number in the fatty acid chain followed by a colon, then the number of double bonds and the position of the double bond closest to the methyl side of the fatty acid molecule. The methyl group is number 1 (the last character in the Greek alphabet is ω, hence the end). In shorthand notation, the unsaturated fatty acids are assumed to have cis bonding and, if the fatty acid is polyunsaturated, double bonds are in the methylene interrupted positions (Fig. 2). In this example, CH₂ (methylene) groups at Δ8 and Δ11 “interrupt” what would otherwise be a conjugated bond system.

Shorthand terminology cannot be used for fatty acids with trans or acetylene bonds, for those with additional functional groups (branched, hydroxy, etc.), or for double-bond systems (≥2 double bonds) that are not methylene interrupted (isolated
### Table 5  Systematic, Common, and Shorthand Names of Unsaturated Fatty Acids

<table>
<thead>
<tr>
<th>Systematic name</th>
<th>Common name</th>
<th>Shorthand</th>
</tr>
</thead>
<tbody>
<tr>
<td>c-9-Dodecenoic</td>
<td>Lauroleic</td>
<td>12:1ω3</td>
</tr>
<tr>
<td>c-5-Tetradecenoic</td>
<td>Physeteric</td>
<td>14:1ω9</td>
</tr>
<tr>
<td>c-9-Tetradecenoic</td>
<td>Myristoleic</td>
<td>14:1ω5</td>
</tr>
<tr>
<td>c-9-Hexadecenoic</td>
<td>Palmitoleic</td>
<td>16:1ω7</td>
</tr>
<tr>
<td>c-7,c-10,c-13-Hexadecatrienoic</td>
<td>—</td>
<td>16:3ω3</td>
</tr>
<tr>
<td>c-4,c-7,c-10,c-13-Hexadecatetraenoic</td>
<td>—</td>
<td>16:4ω3</td>
</tr>
<tr>
<td>c-9-Octadecenoic</td>
<td>Oleic</td>
<td>18:1ω9</td>
</tr>
<tr>
<td>c-11-Octadecenoic</td>
<td>cis-Vaccenic (Asclepic)</td>
<td>18:1ω7</td>
</tr>
<tr>
<td>t-11-Octadecenoic</td>
<td>Vaccenic</td>
<td>*</td>
</tr>
<tr>
<td>t-9-Octadecenoic</td>
<td>Elaidic</td>
<td>*</td>
</tr>
<tr>
<td>c-9,c-12-Octadecadienoic</td>
<td>Linoleic</td>
<td>18:2ω6</td>
</tr>
<tr>
<td>c-9-t-11-Octadecadienoic acid</td>
<td>Ruminic&lt;sup&gt;b&lt;/sup&gt;</td>
<td>*</td>
</tr>
<tr>
<td>c-9,c-12,c-15-Octadecatrienoic</td>
<td>Linolenic</td>
<td>18:3ω3</td>
</tr>
<tr>
<td>c-6,c-9,c-12-Octadecatrienoic</td>
<td>γ-Linolenic</td>
<td>18:3ω6</td>
</tr>
<tr>
<td>c-6,c-9,c-12,c-15-Octadecatetraenoic</td>
<td>Stearidonic</td>
<td>18:4ω3</td>
</tr>
<tr>
<td>c-11-Eicosenoic</td>
<td>Gondoic</td>
<td>20:1ω9</td>
</tr>
<tr>
<td>c-9-Eicosenoic</td>
<td>Gadoic</td>
<td>20:1ω11</td>
</tr>
<tr>
<td>c-8,c-11,c-14-Eicosatrienoic</td>
<td>Dihomo-γ-linolenic</td>
<td>20:3ω6</td>
</tr>
<tr>
<td>c-5,c-8,c-11-Eicosatrienoic</td>
<td>Mead’s</td>
<td>20:3ω9</td>
</tr>
<tr>
<td>c-5,c-8,c-11,c-14-Eicosatrienoic</td>
<td>Arachidonic</td>
<td>20:4ω6</td>
</tr>
<tr>
<td>c-5,c-8,c-11,c-14,c-17-Eicosapentaeno</td>
<td>Eicosapentaeno (EPA)</td>
<td>20:5ω3</td>
</tr>
<tr>
<td>c-13-Docosenoic</td>
<td>Erucic</td>
<td>22:1ω9</td>
</tr>
<tr>
<td>c-11-Docosenoic</td>
<td>Cetoleic</td>
<td>22:1ω11</td>
</tr>
<tr>
<td>c-7,c-10,c-13,c-16,c-19-Docosapentaeno</td>
<td>DPA</td>
<td>22:5ω3</td>
</tr>
<tr>
<td>c-4,c-7,c-10,c-13,c-16,c-19-Docosahexaenoic</td>
<td>DHA</td>
<td>22:6ω3</td>
</tr>
<tr>
<td>c-15-Tetracosenoic</td>
<td>Nervonic (Selacholeic)</td>
<td>24:1ω9</td>
</tr>
</tbody>
</table>

<sup>a</sup>Shorthand nomenclature cannot be used to name trans fatty acids.

<sup>b</sup>One of the conjugated linoleic acid (CLA) isomers.

---

**Figure 2**  IUPAC Δ and common ω numbering systems.
or conjugated). Despite the limitations, shorthand terminology is very popular because of its simplicity and because most of the fatty acids of nutritional importance can be named. Sometimes the ω is replaced by n- (18:2n-6 instead of 18:2ω6). Although there have been recommendations to eliminate ω and use n- exclusively [6], both n- and ω are commonly used in the literature and are equivalent.

Shorthand designations for polyunsaturated fatty acids are sometimes reported without the ω term (18:3). However, this notation is ambiguous, since 18:3 could represent 18:3ω1, 18:3ω3, 18:3ω6, or 18:3ω9; fatty acids, which are completely different in their origins and nutritional significances. Two or more fatty acids with the same carbon and double-bond numbers are possible in many common oils. Therefore, the ω terminology should always be used with the ω term specified.

III. LIPID CLASSES

A. Fatty Acids

1. Saturated Fatty Acids

The saturated fatty acids begin with methanoic (formic) acid. Methanoic, ethanoic, and propanoic acids are uncommon in natural fats and are often omitted from definitions of lipids. However, they are found nonesterified in many food products. Omitting these fatty acids because they are water soluble would argue for also eliminating butyric acid, which would be difficult given its importance in dairy fats. The simplest solution is to accept the very short chain carboxylic acids as fatty acids while acknowledging the rarity in natural fats of these water-soluble compounds. The systematic, common, and shorthand designations of some saturated fatty acids are shown in Table 4.

2. Unsaturated Fatty Acids

By far the most common monounsaturated fatty acid is oleic acid (18:1ω9), although more than 100 monounsaturated fatty acids have been identified in nature. The most common double-bond position for monoenes is ω9. However, certain families of plants have been shown to accumulate what would be considered unusual fatty acid patterns. For example, Eranthis seed oil contains Δ5 monoenes and non-methylene-interrupted polyunsaturated fatty acids containing Δ5 bonds [11]. Erucic acid (22:1ω9) is found at high levels (40–50%) in Cruciferae such as rapeseed and mustard seed. Canola is a rapeseed oil that is low in erucic acid (<3% 22:1ω9).

Polyunsaturated fatty acids (PUFAs) are best described in terms of families because of the metabolism that allows interconversion within, but not among, families of PUFA. The essentiality of ω6 fatty acids has been known since the late 1920s. Signs of ω6 fatty acid deficiency include decreased growth, increased epidermal water loss, impaired wound healing, and impaired reproduction [12,13]. Early studies did not provide clear evidence that ω3 fatty acids are essential. However, since the 1970s, evidence has accumulated illustrating the essentiality of the ω3 PUFA.

Not all PUFAs are EFAs. Plants are able to synthesize de novo and interconvert ω3 and ω6 fatty acid families via desaturases with specificity in the Δ12 and Δ15 positions. Animals have Δ5, Δ6, and Δ9 desaturase enzymes and are unable to synthesized the ω3 and ω6 PUFAs de novo. However, extensive elongation and de-
saturation of EFA occurs (primarily in the liver). The elongation and desaturation of 18:2ω6 is illustrated in Figure 3. The most common of the ω6 fatty acids in our diets is 18:2ω6. Often considered the parent of the ω6 family, 18:2ω6 is first desaturated to 18:3ω6. The rate of this first desaturation is thought to be limiting in premature infants, in the elderly, and under certain disease states. Thus, a great deal of interest has been placed in the few oils that contain 18:3ω6, γ-linolenic acid (GLA). Relatively rich sources of GLA include black currant, evening primrose, and borage oils. GLA is elongated to 20:3ω6, dihomo-γ-linolenic acid (DHGLA). DHGLA is the precursor molecule to the 1-series prostaglandins. DHGLA is further desaturated to 20:4ω6, precursor to the 2-series prostaglandins. Further elongation and desaturation to 22:4ω6 and 22:5ω6 can occur, although the exact function of these fatty acids remains obscure.

Figure 4 illustrates analogous elongation and desaturation of 18:3ω3. The elongation of 20:5ω3 to 22:5ω3 was thought for many years to be via Δ4 desaturase. The inexplicable difficulty in identifying and isolating the putative Δ4 desaturase led to the conclusion that it did not exist, and the pathway from 20:5ω3 to 22:6ω3 was elucidated as a double elongation, desaturation, and β-oxidation.

One of the main functions of the EFAs is their conversion to metabolically active prostaglandins and leukotrienes [14,15]. Examples of some of the possible conversions from 20:4ω6 are shown in Figures 5 and 6 [15]. The prostaglandins are called eicosanoids as a class and originate from the action of cyclooxygenase on 20:4ω6 to produce PGG2. The standard nomenclature of prostaglandins allows usage of the names presented in Figure 5. For a name such as PGG2, the PG represents prostaglandin, the next letter (G) refers to its structure (Fig. 7), and the subscript number refers to the number of double bonds in the molecule.

The parent structure for most of the prostaglandins is prostanoic acid (Fig. 7) [14]. Thus, the prostaglandins can be named based on this parent structure. As well,

---

**Figure 3** Pathway of 18:2ω6 metabolism to 20:4ω6.
they can be named using standard nomenclature rules. For example, prostaglandin E\(_2\) (PGE\(_2\)) is named (5\(Z\),11\(\alpha\),13\(E\),1\(\alpha\))-11,15-dihydroxy-9-oxoprosta-5,13-dienoic acid using the prostanoic acid template. It can also be named using standard nomenclature as 7-[3-hydroxy-2-(3-hydroxy-1-octenyl)-5-oxocyclopentyl]-cis-5-heptenoic acid.

The leukotrienes are produced from 20:4\(\omega 6\) via 5-, 12-, or 15-lipoxygenases to a wide range of metabolically active molecules. The nomenclature is shown in Figure 6.

It is important to realize that there are 1-, 2-, and 3-series prostaglandins originating from 20:3\(\omega 6\), 20:4\(\omega 6\), and 20:5\(\omega 3\), respectively. The structures of the 1- and 3-prostaglandins differ by the removal or addition of the appropriate double bonds. Leukotrienes of the 3-, 4-, and 5-series are formed via lipoxygenase activity on 20:3\(\omega 6\), 20:4\(\omega 6\), and 20:5\(\omega 3\). A great deal of interest has been focused on changing proportions of the prostaglandins and leukotrienes of the various series by diet to modulate various diseases.
Figure 5  Prostaglandin metabolites of 20:4\omega6.
Figure 6  Leucotriene metabolites of 20:4ω6.
3. Acetylenic Fatty Acids

A number of different fatty acids have been identified having triple bonds [16]. The nomenclature is similar to double bonds except that the -ane ending of the parent alkane is replaced with -ynoic acid, -diynoic acid, and so on.

Shorthand nomenclature uses a lowercase $a$ to represent the acetylenic bond; 9c,12a-18:2 is an octadecynoic acid with a double bond in position 9 and the triple bond in position 12. Figure 8 shows the common names and standard nomenclature for some acetylenic fatty acids. Since the ligands attached to triple-bonded carbons are $180^\circ$ from one another (the structure through the bond is linear), the second representation in Figure 8 is more accurate.

The acetylenic fatty acids found in nature are usually 18-carbon molecules with unsaturation starting at $\Delta^9$ consisting of conjugated double–triple bonds [9,16]. Acetylenic fatty acids are rare.

4. trans Fatty Acids

$trans$ Fatty acids include any unsaturated fatty acid that contains double-bond geometry in the $E$ (trans) configuration. Nomenclature differs only from normal $cis$ fatty acids in the configuration of the double bonds.

The three main origins of $trans$ fatty acids in our diet are bacteria, deodorized oils, and partially hydrogenated oils. The preponderance of $trans$ fatty acids in our diets are derived from the hydrogenation process.

Hydrogenation is used to stabilize and improve oxidative stability of oils and to create plastic fats from oils [17]. The isomers that are formed during hydrogenation depend on the nature and amount of catalyst, the extent of hydrogenation, and other factors. The identification of the exact composition of a partially hydrogenated oil is extremely complicated and time consuming. The partial hydrogenation process produces a mixture of positional and geometrical isomers. Identification of the fatty
acid isomers in a hydrogenated menhaden oil has been described [18]. The 20:1 isomers originally present in the unhydrogenated oil were predominantly cis-Δ11 (73% of total 20:1) and cis-Δ13 (15% of total 20:1). After hydrogenation from an initial iodine value of 159 to 96.5, the 20:1 isomers were distributed broadly across the molecules from Δ3 to Δ17 (Fig. 9). The major trans isomers were Δ11 and Δ13, while the main cis isomers were Δ6, Δ9, and Δ11. Similar broad ranges of isomers are produced in hydrogenated vegetable oils [17].

Geometrical isomers of essential fatty acids linoleic and linolenic were first reported in deodorized rapeseed oils [19]. The geometrical isomers that result from deodorization are found in vegetable oils and products made from vegetable oils (infant formulas) and include 9c,12t-18:2, 9t,12c-18:2, and 9t,12r-18:2, as well as 9c,12c,15r-18:3, 9r,12c,15c-18:3, 9c,12r,15c-18:3, and 9t,12c,15r-18:3 [19–22]. These trans-EFA isomers have been shown to have altered biological effects and are incorporated into nervous tissue membranes [23,24], although the importance of these findings has not been elucidated.
trans Fatty acids are formed by some bacteria primarily under anaerobic conditions [25]. It is believed that the formation of trans fatty acids in bacterial cell membranes is an adaptation response to decrease membrane fluidity, perhaps as a reaction to elevated temperature or stress from solvents or other lipophilic compounds that affect membrane fluidity (4-chlorophenol).

Not all bacteria produce appreciable levels of trans fatty acids. The trans-producing bacteria are predominantly gram negative and produce trans fatty acids under anaerobic conditions. The predominant formation of trans is via double-bond migration and isomerization, although some bacteria appear to be capable of isomerization without bond migration. The action of bacteria in the anaerobic rumen results in biohydrogenation of fatty acids and results in trans fatty acid formation in dairy fats (2–6% of total fatty acids). The double bond positions of the trans acids in dairy fats are predominantly in the Δ11 position, with smaller amounts in Δ9, Δ10, Δ13, and Δ14 positions [26].
5. Branched Fatty Acids

A large number of branched fatty acids have been identified [16]. The fatty acids can be named according to rules for branching in hydrocarbons (Table 2). Beside standard nomenclature, several common terms have been retained, including iso-, with a methyl branch on the penultimate (ω2) carbon, and anteiso, with a methyl branch on the antepenultimate (ω3) carbon. The iso and anteiso fatty acids are thought to originate from a modification of the normal de novo biosynthesis, with acetate replaced by 2-methyl propanoate or 2-methylbutanoate, respectively [16]. Other branched fatty acids are derived from isoprenoid biosynthesis including pristanic acid (2,6,10,14-tetramethylpentadecanoic acid) and phytic acid (3,7,11,15-tetramethylhexadecanoic acid).

6. Cyclic Fatty Acids

Many fatty acids that exist in nature contain cyclic carbon rings [27]. Ring structures contain either three (cyclopropyl and cyclopropenyl), five (cyclopentenyl), or six (cyclohexenyl) carbon atoms and may be saturated or unsaturated. As well, cyclic fatty acid structures resulting from heating the vegetable oils have been identified [27–29].

In nomenclature of cyclic fatty acids, the parent fatty acid is the chain from the carboxyl group to the ring structure. The ring structure and additional ligands are considered a substituent of the parent fatty acid. An example is given in Figure 10. The parent in this example is nonanoic acid (not pentadecanoic acid, which would result if the chain were extended through the ring structure). The substituted group is a cyclopentyl group with a 2-butyl ligand (2-butylcyclopentyl). Thus the correct standard nomenclature is 9-(2-butylcyclopentyl)nonanoic acid. The 2 is sometimes expressed as 2’ to indicate that the numbering is for the ring, and not the parent chain. The C-1 and C-2 carbons of the cyclopentyl ring are chiral, and two possible configurations are possible. Both the carboxyl and longest hydrocarbon substituents can be on the same side of the ring, or they can be on opposite sides. These are referred to as cis and trans, respectively.

The cyclopropene and cyclopropane fatty acids can be named by means of the standard nomenclature noted in the example above. They are also commonly named using the parent structure that carries through the ring structure. In the example in Figure 11, the fatty acid (commonly named lactobacillic acid or phycomonic acid) is named 10-(2-hexylcyclopropyl)decanonic acid in standard nomenclature. An older naming system would refer to this fatty acid as cis-11,12-methyleneoctadecanoic acid, where cis designates the configuration of the ring structure. If the fatty acid is unsaturated, the term methylene is retained but the double bond position is noted in the parent fatty acid structure (cis-11,12-methylene-cis-octadec-9-enoic acid).

Figure 12 presents some examples of natural cyclic fatty acids and their trivial and standard nomenclature.

7. Hydroxy and Epoxy Fatty Acids

Saturated and unsaturated fatty acids containing hydroxy and epoxy functional groups have been identified [1,16]. Hydroxy fatty acids are named by means of the parent fatty acid and the hydroxy group(s) numbered with its Δ location. For example, the fatty acid with the trivial name ricinoleic (Fig. 13) is named R-12-hy-
droxy-cis-9-octadecenoic acid. Ricinoleic acid is found in the seeds of *Ricinus* species and accounts for about 90% of the fatty acids in castor bean oil.

Because the hydroxy group is chiral, stereoisomers are possible. The *R/S* system is used to identify the exact structure of the fatty acid. Table 6 reviews the rules for *R/S* nomenclature. The *R/S* system can be used instead of the *α/β* and *cis/trans* nomenclature systems. A fatty acid with a hydroxy substituent in the Δ2 position is commonly called an *α*-hydroxy acid; fatty acids with hydroxy substituents in the Δ3 and Δ4 positions are called *β*-hydroxy acids and *γ*-hydroxy acids, respectively. Some

**Figure 10** Nomenclature of cyclic fatty acids.

cis-11,12-methyleneoctadecanoic acid

cis-10-(2-hexylcyclopropyl)decanoic acid

**Figure 11** Nomenclature for a cyclopropenoid fatty acid.
Cyclic fatty acid structures and nomenclature.

Figure 12  Cyclic fatty acid structures and nomenclature.

common hydroxy acids are shown in Figure 13. Cutins, which are found in the outer layer of fruit skins, are composed of hydroxy acid polymers, which also may contain epoxy groups [16].

Epoxy acids, found in some seed oils, are formed on prolonged storage of seeds [16]. They are named similarly to cyclopropane fatty acids, with the parent acid considered to have a substituted oxirane substituent. An example of epoxy fatty acids and their nomenclature is shown in Figure 14. The fatty acid with the common name vernolic acid is named (using standard nomenclature) 11-(3-pentyloxyranyl)-9-undecenoic acid. In older nomenclature, where the carbon chain is carried through the oxirane ring, vernolic acid would be called 12,13-epoxyoleic acid or 12-13-epoxy-9-octadecenoic acid. The configuration of the oxirane ring substituents can be named in the cis/trans, E/Z, or R/S configuration systems.
8. Furanoid Fatty Acids

Some fatty acids contain an unsaturated oxolane heterocyclic group. There are more commonly called furanoid fatty acids because a furan structure (diunsaturated oxolane) is present in the molecule. Furanoid fatty acids have been identified in *Exocarpus* seed oils. They have also been identified in plants, algae, and bacteria and are a major component in triacylglycerols from latex rubber [1,16]. They are important in marine oils and may total several percentage points of the total fatty acids or more in liver and testes [1,30].

Furanoid fatty acids have a general structure shown in Figure 15. A common nomenclature describing the furanoid fatty acids (as F₁, F₂, etc.) is used [30]. The naming of the fatty acids in this nomenclature is arbitrary and originated from elution order in gas chromatography. A shorthand notation that is more descriptive gives the methyl substitution followed by F, and then the carbon lengths of the carboxyl and

---

**Table 6**  A Summary of Rules for *R/S* Nomenclature

<table>
<thead>
<tr>
<th>Rule</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>The sequence priority rules (Table 3) are used to prioritize the ligands attached to the chiral center ((a &gt; b &gt; c &gt; d)).</td>
</tr>
<tr>
<td>2.</td>
<td>The molecule is viewed with the (d) substituent facing away from the viewer.</td>
</tr>
<tr>
<td>3.</td>
<td>The remaining three ligands ((a, b, c)) will be oriented with the order (a-b-c) in a clockwise or counterclockwise direction.</td>
</tr>
<tr>
<td>4.</td>
<td>Clockwise describes the (R) (rectus, right) conformation, and counterclockwise describes the (S) (sinister, left) conformation.</td>
</tr>
</tbody>
</table>

*Source: Ref. 10.*
terminal chains in parentheses: MeF(9,5). Standard nomenclature follows the same principles outlined in Sec. IV.A.6. The parent fatty acid chain extends only to the furan structure, which is named as a ligand attached to the parent molecule. For example, the fatty acid named F5 in Figure 15 is named 11-(3,4-dimethyl-5-pentyl-2-furyl)undecanoic acid. Shorthand notation for this fatty acid would be F5 or MeF(11,5). Numbering for the furan ring starts at the oxygen and proceeds clockwise.

B. Acylglycerols

Acylglycerols are the predominant constituent in oils and fats of commercial importance. Glycerol can be esterified with one, two, or three fatty acids, and the individual fatty acids can be located on different carbons of glycerol. The terms monoacylglycerol, diacylglycerol, and triacylglycerol are preferred for these compounds over the older and confusing names mono-, di-, and triglycerides [6,7].

Fatty acids can be esterified on the primary or secondary hydroxyl groups of glycerol. Although glycerol itself has no chiral center, it becomes chiral if different fatty acids are esterified to the primary hydroxyls or if one of the primary hydroxyls
Figure 15  Furanoid fatty acid structure and shorthand nomenclature.

<table>
<thead>
<tr>
<th>Name</th>
<th>x</th>
<th>y</th>
<th>R</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>8</td>
<td>2</td>
<td>CH₃</td>
</tr>
<tr>
<td>F2</td>
<td>8</td>
<td>4</td>
<td>H</td>
</tr>
<tr>
<td>F3</td>
<td>8</td>
<td>4</td>
<td>CH₃</td>
</tr>
<tr>
<td>F4</td>
<td>10</td>
<td>2</td>
<td>CH₃</td>
</tr>
<tr>
<td>F5</td>
<td>10</td>
<td>4</td>
<td>H</td>
</tr>
<tr>
<td>F6</td>
<td>10</td>
<td>4</td>
<td>CH₃</td>
</tr>
<tr>
<td>F7</td>
<td>12</td>
<td>4</td>
<td>H</td>
</tr>
<tr>
<td>F8</td>
<td>12</td>
<td>4</td>
<td>CH₃</td>
</tr>
</tbody>
</table>

Figure 16  Chiral carbons in acylglycerols.

is esterified. Thus, terminology must differentiate between the two possible configurations (Fig. 16). The most common convention to differentiate these stereoisomers is the sn convention of Hirshmann (see Ref. 31). In the numbering that describes the hydroxyl groups on the glycerol molecule in Fisher projection, sn1, sn2, and sn3 designations are used for the top (C1), middle (C2), and bottom (C3) OH groups (Fig. 17). The sn term indicates stereospecific numbering [1].

C* = chiral carbon
In common nomenclature, esters are called α on primary and β on secondary OH groups. If the two primary-bonded fatty acids are present, the primary carbons are called α and α'. If one or two acyl groups are present, the term “partial glyceride” is sometimes used. Nomenclature of the common partial glycerides is shown in Figure 18.

Standard nomenclature allows several different names for each triacylglycerol (TAG) [6]. A TAG with three stearic acid esters can be named as glycerol tristearate, tristearoyl glycerol, or tri-Ω-stearoyl glycerol. The “Ω” locant can be omitted if the fatty acid is esterified to the hydroxyl group. More commonly, triacylglycerol nomenclature uses the designation -in to indicate the molecule in a TAG (e.g., tristearin). If different fatty acids are esterified to the TAG—for example, the TAG with sn-1 palmitic acid, sn-2 oleic acid, and sn-3 stearic acid—the name replaces the -ic in the fatty acid name with -oyl, and fatty acids are named in sn1, sn2, sn3 order (1-palmitoyl-2-oleoyl-3-stearoyl-sn-glycerol). This TAG also can be named as sn-1-palmito-2-oleo-3-stearin or sn-glycerol-1-palmate-2-oleate-3-stearate. If two of the fatty acids are identical, the name incorporates the designation di- (e.g., 1,2-dipalmitoyl-3-oleoyl-sn-glycerol, 1-stearoyl-2,3-dilinolenoyl-sn-glycerol, etc.).
Table 7  Short Abbreviations for Some Common Fatty Acids

<table>
<thead>
<tr>
<th>AC</th>
<th>Acetic</th>
<th>Ln</th>
<th>Linolenic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ad</td>
<td>Arachidic</td>
<td>M</td>
<td>Myristic</td>
</tr>
<tr>
<td>An</td>
<td>Arachidonic</td>
<td>N</td>
<td>Nervonic</td>
</tr>
<tr>
<td>B</td>
<td>Butyric</td>
<td>O</td>
<td>Oleic</td>
</tr>
<tr>
<td>Be</td>
<td>Behenic</td>
<td>Oc</td>
<td>Octanoic</td>
</tr>
<tr>
<td>D</td>
<td>Decanoic</td>
<td>P</td>
<td>Palmitic</td>
</tr>
<tr>
<td>E</td>
<td>Erucic</td>
<td>Po</td>
<td>Palmitoleic</td>
</tr>
<tr>
<td>El</td>
<td>Elaidic</td>
<td>R</td>
<td>Ricinoleic</td>
</tr>
<tr>
<td>G</td>
<td>Eicosenoic</td>
<td>S</td>
<td>Saturated (any)</td>
</tr>
<tr>
<td>H</td>
<td>Hexanoic</td>
<td>St</td>
<td>Stearic</td>
</tr>
<tr>
<td>L</td>
<td>Linoleic</td>
<td>U</td>
<td>Unsaturated (any)</td>
</tr>
<tr>
<td>La</td>
<td>Lauric</td>
<td>V</td>
<td>Vaccenic</td>
</tr>
<tr>
<td>Lg</td>
<td>Lignoceric</td>
<td>X</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

Source: Ref. 31.

To facilitate TAG descriptions, fatty acids are abbreviated using one or two letters (Table 7). The triacylglycerols can be named after the esterified fatty acids using shorthand nomenclature. For example, sn-POSt is shorthand description for the molecule 1-palmitoyl-2-oleoyl-3-stearoyl-sn-glycerol. If the sn- is omitted, the stereospecific positions of the fatty acids are unknown. POSt could be a mixture of sn-POSt, sn-StOP, sn-PStO, sn-OPSt, or sn-StPO, in any proportion. An equal mixture of both stereoisomers (the racemate) is designated as rac. Thus,rac-OPP represents equal amounts of sn-OPP and sn-PPO. If only the sn-2 substituent is known with certainty in a TAG, the designation β- is used. For example, β-POSt is a mixture (unknown amounts) of sn-POSt and sn-StOP.

TAGs are also sometimes described by means of the ω nomenclature. For example, sn-18:0-18:2/ω6-16:0 represents 1-stearoyl-2-linoleoyl-3-palmitoyl-sn-glycerol.

C. Sterols and Sterol Esters

The steroid class of organic compounds includes sterols of importance in lipid chemistry. Although the term “sterol” is widely used, it has never been formally defined. The following working definition was proposed some years ago: “Any hydroxylated steroid that retains some or all of the carbon atoms of squalene in its side chain and partitions nearly completely into the ether layer when it is shaken with equal volumes of ether and water” [32]. Thus, for this definition, sterols are a subset of steroids and exclude the steroid hormones and bile acids. The importance of bile acids and their intimate origin from cholesterol makes this definition difficult. As well, non-hydroxylated structures such as cholestane, which retain the steroid structure, are commonly considered sterols.

The sterols may be derived from plant (phytosterols) or animal (zoosterols) sources. They are widely distributed and are important in cell membranes. The predominant zoosterol is cholesterol. Although a few phytosterols predominate, the sterol composition of plants can be very complex. For example, as many as 65 different sterols have been identified in corn (Zea mays) [33].
In the standard ring and carbon numbering (Fig. 19) [33], the actual three-dimensional configuration of the tetra ring structure is almost flat, so the ring substituents are either in the same plane as the rings or in front or behind the rings. If the structure in Figure 19 lacks one or more of the carbon atoms, the numbering of the remainder will not be changed.

The methyl group at position 10 is axial and lies in front of the general plane of the molecule. This is the \( \beta \) configuration and is designated by connection using a solid or thickened line. Atoms or groups behind the molecule plane are joined to the ring structure by a dotted or broken line and are given the \( \alpha \) configuration. If the stereochemical configuration is not known, a wavy line is used and the configuration is referred to as \( \epsilon \). Unfortunately, actual three-dimensional position of the substituents may be in plane, in front of, or behind the plane of the molecule. The difficulties with this nomenclature have been discussed elsewhere [32,33].

The nomenclature of the steroids is based on parent ring structures. Some of the basic steroid structures are presented in Figure 20 [6]. Because cholesterol is a derivative of the cholestane structure (with the H at C-5 eliminated because of the double bond), the correct standard nomenclature for cholesterol is 3\( \beta \)-cholest-5-en-3-ol. The complexity of standardized nomenclature has led to the retention of trivial names for some of the common structures (e.g., cholesterol). However, when the structure is changed—for example, with the addition of a ketone group to cholesterol at the 7-position—the proper name is 3\( \beta \)-hydroxycholest-5-en-7-one, although this molecule is also called 7-ketocholesterol in common usage.

A number of other sterols of importance in foods are shown in Figure 21. The trivial names are retained for these compounds, but based on the nomenclature system discussed for sterols, stigmasterol can be named 3\( \beta \)-hydroxy-24-ethylcholesta-5,22-diene. Recent studies have suggested that plant sterols and stanols (saturated derivatives of sterols) have cholesterol lowering properties in humans [33a].

Cholesterol has been reported to oxidize in vivo and during food processing [34–37]. These cholesterol oxides have come under intense scrutiny because they have been implicated in development of atherosclerosis. Some of the more commonly

---

**Figure 19** Carbon numbering in cholesterol structure.
reported oxidation products are shown in Figures 22 and 23. Nomenclature in common usage in this field often refers to the oxides as derivatives of the cholesterol parent molecule: 7-β-hydroxycholesterol, 7-ketocholesterol, 5,6β-epoxycholesterol, and so on. The standard nomenclature follows described rules and is shown in Figures 22 and 23.
Sterol esters exist commonly and are named using standard rules for esters. For example, the ester of cholesterol with palmitic acid would be named cholesterol palmitate. The standard nomenclature would also allow this molecule to be named 3-O-palmitoyl-3β-cholest-5-en-3-ol or 3-palmitoyl-3β-cholest-5-en-3-ol.

D. Waxes

Waxes (commonly called wax esters) are esters of fatty acids and long chain alcohols. Simple waxes are esters of medium chain fatty acids (16:0, 18:0, 18:1ω9) and long chain aliphatic alcohols. The alcohols range in size from C8 to C18. Simple waxes are found on the surface of animals, plants, and insects and play a role in prevention
Figure 22  Cholesterol oxidation products and nomenclature I. (From Ref. 37.)
Figure 23  Cholesterol oxidation products and nomenclature II. (From Ref. 37.)

of water loss. Complex waxes are formed from diols or from alcohol acids. Di- and triesters as well as acid and alcohol esters have been described. Simple waxes can be named by removing the -ol from the alcohol and replacing it with -yl, and replacing the -ic from the acid with -oate. For example, the wax ester from hexadecanol and oleic acid would be named hexadecyl oleate or hexadecyl-cis-9-hexadenedenoate. Some of the long chain alcohols have common names derived from the fatty acid parent (e.g., lauryl alcohol, stearyl alcohol). The C16 alcohol (1-hexadecanol) is commonly called cetyl alcohol. Thus, cetyl oleate is another acceptable name for this compound.

Waxes are found in animal, insect, and plant secretions as protective coatings. Waxes of importance in foods as additives include beeswax, carnauba wax, and candelilla wax.
E. Phosphoglycerides (Phospholipids)

Phosphoglycerides (PLs) are composed of glycerol, fatty acids, phosphate, and (usually) an organic base or polyhydroxy compound. The phosphate is almost always linked to the \( sn-3 \) position of glycerol molecule.

The parent structure of the phosphoglycerides is phosphatidic acid (\( sn-1,2\)-diacylglycerol-3-phosphate). The terminology for phosphoglycerides is analogous to that of acylglycerols with the exception of the no acyl group at \( sn-3 \). The prefix _lyso-,_ when used for phosphoglycerides, indicates that the \( sn-2 \) position has been hydrolyzed and a fatty acid is esterified to the \( sn-1 \) position only.

Some common phosphoglyceride structures and nomenclature are presented in Figure 24. Phospholipid classes are denoted using shorthand designation (PC = phosphatidylcholine, etc.). The standard nomenclature is based on the PL type. For example, a PC with an oleic acid on \( sn-1 \) and linolenic acid on \( sn-2 \) would be named 1-oleoyl-2-linolenoyl-\( sn \)-glycerol-3-phosphocholine. The name phosphorycholine is sometimes used but is not recommended [8]. The terms lecithin and cephalin, sometimes used for PC and PE, respectively, are not recommended [8].

\[
\begin{align*}
\text{E. Phosphoglycerides (Phospholipids)}
\end{align*}
\]

\[
\begin{align*}
\text{Figure 24} \quad \text{Nomenclature for glycerophospholipids.}
\end{align*}
\]
Cardiolipin is a phosphoglyceride that is present in heart muscle mitochondria and bacterial membranes. Its structure and nomenclature are shown in Figure 25. Some cardiolipins contain the maximum possible number of 18:2 molecules (4 mol/mol).

F. Ether(phospho)glycerides (Plasmalogens)

Plasmalogens are formed when a vinyl (1-alkenyl) ether bond is found in a phospholipid or acylglycerol. The 1-alkenyl-2,3-diacylglycerols are termed neutral plasmalogens. A 2-acyl-1-(1-alkenyl)-sn-glycerophosphocholine is named a plasmalogen or plasmenylcholine. The related 1-alkyl compound is named plasmanylcholine.

G. Glyceroglycolipids (Glycosylglycolipids)

The glyceroglycolipids or glycolipids are formed when a 1,2-diacyl-sn-3-glycerol is linked via the sn-3 position to a carbohydrate molecule. The carbohydrate is usually a mono- or a disaccharide, less commonly a tri- or tetrasaccharide. Galactose is the most common carbohydrate molecule in plant glyceroglycolipids. Structures and nomenclature for some glyceroglycolipids are shown in Figure 26. The names monogalactosyldiacylglycerol (MGDG) and digalactosyldiacylglycerol (DGDG) are used in common nomenclature. The standard nomenclature identifies the ring structure and bonding of the carbohydrate groups (Fig. 26).

H. Sphingolipids

The glycosphingolipids are a class of lipids containing a long chain base, fatty acids, and various other compounds, such as phosphate and monosaccharides. The base is commonly sphingosine, although more than 50 bases have been identified. The ceramides are composed of sphingosine and a fatty acid (Fig. 27). Sphingomyelin is one example of a sphingophospholipid. It is a ceramide with a phosphocholine group connected to the primary hydroxyl of sphingosine. The ceramides can also be attached to carbohydrate molecules (sphingoglycolipids or cerebrosides) via the primary hydroxyl group of sphingosine. Gangliosides are complex cerebrosides with the ceramide residue connected to a carbohydrate containing glucose-galactosamine-N-acetyleneuraminic acid. These lipids are important in cell membranes and the brain, and they act as antigenic sites on cell surfaces. Nomenclature and structures of some cerebrosides are shown in Figure 27.
I. Fat-Soluble Vitamins

1. Vitamin A

Vitamin A exists in the diet in many forms (Fig. 28). The most bioactive form is the all-trans retinol, and cis forms are created via light-induced isomerization (Table 8). The 13-cis isomer is the most biopotent of the mono- and di-cis isomers. The α- and β-carotenes have biopotencies of about 8.7% and 16.7% of the all-trans retinol activity, respectively. The daily value (DV) for vitamin A is 1000 retinol equivalents (RE), which represents 1000 μg of all-trans retinol or 6000 μg of β-carotene. Vitamin A can be toxic when taken in levels exceeding the %DV. Some reports suggest that levels of 15,000 RE per day can be toxic [38].
Figure 27  Sphingolipid structures and nomenclature.
**Figure 28** Structures of some vitamin A compounds.

**Table 8** Approximate Biological Activity Relationships of Vitamin A Compounds

<table>
<thead>
<tr>
<th>Compound</th>
<th>Activity of all-trans retinol (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All-trans retinol</td>
<td>100</td>
</tr>
<tr>
<td>9-cis Retinol</td>
<td>21</td>
</tr>
<tr>
<td>11-cis Retinol</td>
<td>24</td>
</tr>
<tr>
<td>13-cis Retinol</td>
<td>75</td>
</tr>
<tr>
<td>9,13-di-cis Retinol</td>
<td>24</td>
</tr>
<tr>
<td>11,13-di-cis Retinol</td>
<td>15</td>
</tr>
<tr>
<td>α-Carotene</td>
<td>8.4</td>
</tr>
<tr>
<td>β-Carotene</td>
<td>16.7</td>
</tr>
</tbody>
</table>
Toxic symptoms of hypervitaminosis A include drowsiness, headache, vomiting, and muscle pain. Vitamin A can be teratogenic at high doses [38]. Vitamin A deficiency results in night blindness and ultimately total blindness, abnormal bone growth, increased cerebrospinal pressure, reproductive defects, abnormal cornification, loss of mucus secretion cells in the intestine, and decreased growth. The importance of beef liver, an excellent source of vitamin A, in cure of night blindness was known to the ancient Egyptians about 1500 BC [39].

2. Vitamin D

Although as many as five vitamin D compounds have been described (Fig. 29), only two of these are biologically active: ergocalciferol (vitamin D₂) and cholecalciferol (vitamin D₃). Vitamin D₃ can be synthesized in humans from 7-dehydrocholesterol, which occurs naturally in the skin via light irradiation (Fig. 30).

The actual hormonal forms of the D vitamins are the hydroxylated derivatives. Vitamin D is converted to 25-OH-D in the kidney and further hydroxylated to 1,25-diOH-D in the liver. The dihydroxy form is the most biologically active form in humans.

3. Vitamin E

Vitamin E compounds include the tocopherols and tocotrienols. Tocotrienols have a conjugated triene double bond system in the phytyl side chain, while tocopherols do not. The basic nomenclature is shown in Figure 31. The bioactivity of the various vitamin E compounds is shown in Table 9. Methyl substitution affects the bioactivity of vitamin E, as well as its in vitro antioxidant activity.

4. Vitamin K

Several forms of vitamin K have been described (Fig. 32). Vitamin K₁ (phylloquinone) is found in green leaves and vitamin K₂ (menaquinone) is synthesized by intestinal bacteria. Vitamin K is involved in blood clotting as an essential cofactor in the synthesis of γ-carboxyglutamate necessary for active prothrombin. Vitamin K deficiency is rare because of intestinal microflora synthesis. Warfarin and dicoumerol prevent vitamin K regeneration and may result in fatal hemorrhaging.

J. Hydrocarbons

The hydrocarbons include normal, branched, saturated, and unsaturated compounds of varying chain lengths. The nomenclature for hydrocarbons has already been discussed. The hydrocarbons of most interest to lipid chemists are the isoprenoids and their oxygenated derivatives.

The basic isoprene unit (2-methyl-1,3-butadiene) is the building block for a large number of interesting compounds, including carotenoids (Fig. 33), oxygenated carotenoids (Fig. 34), sterols, and unsaturated and saturated isoprenoids (isopranes). Recently, it has been discovered that 15-carbon and 20-carbon isoprenoids are covalently attached to some proteins and may be involved in control of cell growth [40]. Members of this class of protein-isoprenoid molecules are called prenylated proteins.

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Figure 29  Structures of some vitamin D compounds.

Figure 30  Formation of vitamin D in vivo.
Figure 31 Structures of some vitamin E compounds.

Table 9 Approximate Biological Activity Relationships of Vitamin E Compounds

<table>
<thead>
<tr>
<th>Compound</th>
<th>Activity of $d$-(\alpha)-tocopherol (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$d$-(\alpha)-Tocopherol</td>
<td>100</td>
</tr>
<tr>
<td>$l$-(\alpha)-Tocopherol</td>
<td>26</td>
</tr>
<tr>
<td>DL-(\alpha)-Tocopherol</td>
<td>74</td>
</tr>
<tr>
<td>DL-(\alpha)-Tocopheryl acetate</td>
<td>68</td>
</tr>
<tr>
<td>$d$-(\beta)$\rightarrow$-Tocopherol</td>
<td>8</td>
</tr>
<tr>
<td>$d$-(\gamma)$\rightarrow$-Tocopherol</td>
<td>3</td>
</tr>
<tr>
<td>$d$-(\delta)$\rightarrow$-Tocopherol</td>
<td>—</td>
</tr>
<tr>
<td>$d$-(\alpha)$\rightarrow$-Tocotrienol</td>
<td>22</td>
</tr>
<tr>
<td>$d$-(\beta)$\rightarrow$-Tocotrienol</td>
<td>3</td>
</tr>
<tr>
<td>$d$-(\gamma)$\rightarrow$-Tocotrienol</td>
<td>—</td>
</tr>
<tr>
<td>$d$-(\delta)$\rightarrow$-Tocotrienol</td>
<td>—</td>
</tr>
</tbody>
</table>
Figure 32  Structures of some vitamin K compounds.

Figure 33  Structures and nomenclature of carotenoids.
Figure 34  Structures and nomenclature of some oxygenated carotenoids.

IV. SUMMARY

It would be impossible to describe the structures and nomenclature of all known lipids even in one entire book. The information presented in this chapter is a brief overview of the complex and interesting compounds we call lipids.

REFERENCES


Chemistry and Function of Phospholipids

Marilyn C. Erickson

The University of Georgia, Griffin, Georgia

I. INTRODUCTION

Phospholipids can generally be regarded as asymmetrical phosphoric acid diesters comprising chemical bonds of three types: C-C bonds, ester bonds, and phosphoester bonds. While hydrolysis is inherent to the ester and phosphoester bonds, other physical and chemical reactions associated with phospholipids are dictated by the kind of head group and by the chain length and degree of unsaturation of the constituent aliphatic moieties. These activities constitute the focus of this chapter. In addition, the ramifications of phospholipids’ amphiphilic nature and their propensity to aggregate as bilayers will be discussed in relation to their functional role in foods.

II. PHOSPHOLIPID CLASSIFICATION

Phospholipids are divided into two main classes depending on whether they contain a glycerol or a sphingosyl backbone (Fig. 1). These differences in base structure affect their chemical reactivity.

Glycerophospholipids are named after and contain structures that are based on phosphatidic acid. The moiety attached to the phosphate includes nitrogenous bases or polyols. Sphingolipids are lipids that contain sphingosine (trans-d-erythro-1,3-dihydroxy-2-amino-4-octadecene) or a related amino alcohol. Although the most common sphingophospholipid, sphingomyelin, represents a major lipid in certain membranes of animals, it is of minor importance in plants and probably is absent from bacteria.
Figure 1  Structure of phospholipids. Circled areas show distinguishing features of each phospholipid.

In most tissues the diacyl forms of the glycerophospholipids predominate, but small amounts of ether derivatives are also found. These are the monoacyl monoalk-1-enyl ether forms of phospholipids. Choline and ethanolamine plasmalogens are the most common forms, although serine plasmalogen has also been found.

The phosphonolipids that contain a covalent bond between the phosphorus atom and the carbon of the nitrogenous base comprise another glycerophospholipid.
variant [1]. Phosphonolipids are major constituents in three phyla and are synthesized by phytoplankton, the base of the food chains of the ocean.

III. PHOSPHOLIPID MESOPHASES

Phospholipids are characterized by the presence of a polar or hydrophilic head group and a nonpolar or hydrophobic fatty acid region. It is this amphipathic character that drives the macroassembly of phospholipids in the presence of water to a bilayer organization in which the polar regions tend to orient toward the aqueous phase and the hydrophobic regions are sequestered from water (Fig. 2A). Another macromolecular structure commonly adopted by phospholipids and compatible with their amphipathic constraints is the hexagonal (HII) phase (Fig. 2B). This phase consists of a hydrocarbon matrix penetrated by hexagonally packed aqueous cylinders with diameters of about 20 Å. Table 1 lists less common macromolecular structures that may be adopted by phospholipids in a solid or liquid state. The ability of phospholipids to adopt these different structures is referred to as lipid polymorphism. Additional information on structure and properties of these mesophases of phospholipids may be found in the review of Seddon and Cevc [2].

IV. BIOLOGICAL MEMBRANES

Phospholipids, along with proteins, are major components of biological membranes, which in turn are an integral part of prokaryotes (bacteria) and eukaryotes (plants and animals). The predominant structures assumed by phospholipids in membranes are the bilayer and HII structure, which is dictated by the phase preference of the individual phospholipids (Table 2). It is immediately apparent that a significant proportion of membrane lipids adopt or promote HII phase structure under appropriate conditions. The most striking example is phosphatidylethanolamine (PE), which may compose up to 30% of membrane phospholipids. Under such conditions, portions of the membrane that adopt an HII phase would be expected to be incompatible with maintenance of a permeability barrier between external and internal compartments at those areas. Consequently, alternative roles for those structures must exist.

A. Membrane Permeability

The ability of lipids to provide a bilayer permeability barrier between external and internal environments constitutes one of their most important functions in a biological

![Figure 2 Mesomorphic structures of phospholipids: (A) lamellar and (B) hexagonal II.](image)
membrane. Permeability coefficients of liquid crystalline lipid bilayers for water are in the range of $10^{-2}$–$10^{-4}$ cm/s, indicating a high permeability [3]. The relative permeability of different membrane systems to water can be monitored by means of light scattering techniques that measure swelling rates when osmotic gradients are applied [4]. Results obtained from such studies indicate that increased unsaturation of the fatty acids of the membrane causes increases in water permeability. Since cholesterol reduces water permeability, the general conclusion has been made that factors contributing to increased order in the hydrocarbon region reduce water permeability.

The diffusion properties of nonelectrolytes (uncharged polar solutes) also appears to depend on the properties of the lipid matrix in much the same manner as does the diffusion of water. That is, decreased unsaturation of phospholipids or increased cholesterol content results in lower permeability coefficients. In the case of

<table>
<thead>
<tr>
<th>Phase</th>
<th>Phase structure</th>
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<tbody>
<tr>
<td>Liquid</td>
<td>Fluid lamellar</td>
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<td></td>
<td>Hexagonal</td>
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<td>Complex hexagonal</td>
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<td>Rhombohedral</td>
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<td>Solid</td>
<td>Three-dimensional crystal</td>
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<td>Rippled gel</td>
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<td>Partial gel</td>
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<tr>
<th>Table 1</th>
<th>Macromolecular Mesophases Adopted by Phospholipids</th>
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<tr>
<td>Phase</td>
<td>Phase structure</td>
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<tr>
<td>Liquid</td>
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<td>Solid</td>
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<td>Partial gel</td>
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<tr>
<th>Table 2</th>
<th>Phase Preference of Membrane Phospholipids</th>
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<tbody>
<tr>
<td>Bilayer</td>
<td>Hexagonal $H_{II}$</td>
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<td>-----------------------------------------------------</td>
</tr>
<tr>
<td>Phosphatidylcholine</td>
<td>Phosphatidylethanolamine</td>
</tr>
<tr>
<td>Sphingomyelin</td>
<td>Phosphatidylserine (pH &lt; 3)</td>
</tr>
<tr>
<td>Phosphatidylserine</td>
<td>Phosphatidylglycerol</td>
</tr>
<tr>
<td>Phosphatidylinositol</td>
<td>Phosphatidic acid (+ Ca$^{2+}$)</td>
</tr>
<tr>
<td>Phosphatidic acid</td>
<td>Phosphatidic acid pH &lt; 3</td>
</tr>
</tbody>
</table>
nonelectrolytes, however, the permeability coefficients are at least two orders of magnitude smaller than those of water. Furthermore, for a given homologous series of compounds, the permeability increases as the solubility in a hydrocarbon environment increases, indicating that the rate-limiting step in diffusion is the initial partitioning of the molecule into the lipid bilayer [5].

Measures of the permeability of membranes to small ions are complicated, since for free permeation to proceed, a counterflow of other ions of equivalent charge is required. In the absence of such a counterflow, a membrane potential is established that is equal and opposite to the chemical potential of the diffusing species. A remarkable impermeability of lipid bilayers exists for small ions with permeability coefficients of less than $10^{-10}$ cm/s commonly observed. While permeability coefficients for Na$^+$ and K$^+$ may be as small as $10^{-14}$ cm/s, lipid bilayers appear to be much more permeable to H$^+$ or OH$^-$ ions, which have been reported to have permeability coefficients in the range of $10^{-4}$ cm/s [6]. One of the hypotheses put forth to explain this anomaly involves hydrogen-bonded wires across membranes. Such water wires could have transient existence in lipid membranes, and when such structures connect the two aqueous phases, proton flux could result as a consequence of H–O–H → O–H bond rearrangements. Such a mechanism does not involve physical movement of a proton all the way across the membrane; hence, proton flux occurring by this mechanism is expected to be significantly faster when compared with the flux of other monovalent ions which lack such a mechanism. As support for the existence of this mechanism, an increase in the level of cholesterol decreased the rate of proton transport that correlated to the decrease in the membrane’s water content [7].

Two alternative mechanisms are frequently used to describe ionic permeation of lipid bilayers. In the first, the solubility-diffusion mechanism, ions partition and diffuse across the hydrophobic phase. In the second, the pore mechanism, ions traverse the bilayer through transient hydrophilic defects caused by thermal fluctuations. Based on the dependence of halide permeability coefficients on bilayer thickness and on ionic size, a solubility-diffusion mechanism was ascribed to these ions [8]. In contrast, permeation by monovalent cations, such as potassium, has been accounted for by a combination of both mechanisms. In terms of the relationship between lipid composition and membrane permeability, ion permeability appears to be related to the order in the hydrocarbon region, where increased order leads to a decrease in permeability. The charge on the phospholipid polar head group can also strongly influence permeability by virtue of the resulting surface potential. Depending on whether the surface potential is positive or negative, anions and cations could be attracted or repelled to the lipid–water interface.

**B. Membrane Fluidity**

The current concept of biological membranes is a dynamic molecular assembly characterized by the coexistence of structures with highly restricted mobility and components having great rotational freedom. These membrane lipids and proteins comprising domains of highly restricted mobility appear to exist on a micrometer scale in a number of cell types [9,10]. Despite this heterogeneity, membrane fluidity is still considered as a bulk, uniform property of the lipid phase that is governed by a complex pattern of the components’ mobilities. Individual lipid molecules can display
diffusion of three different types: lateral, rotational, and transversal [11]. Lateral diffusion of lipids in biological membranes refers to the two-dimensional translocation of the molecules in the plane of the membrane. Rotational diffusion of lipid molecules is restricted to the plane of biological membranes, whereas transverse diffusion is the out-of-plane rotation or redistribution of lipid molecules between the two leaflets of the bilayer. Transverse diffusion or “flip-flop” motion is very low in lipid bilayers, and specific enzymes are required to mediate the process.

There are two major components of membrane fluidity. The first component is the order parameter ($S$), also called the structural, static, or range component of membrane fluidity. This is a measure of angular range of rotational motion, with more tightly packed chains resulting in a more ordered or less fluid bilayer. The second component of membrane fluidity is microviscosity and is the dynamic component of membrane fluidity. This component measures the rate of rotational motion and is a more accurate reflection of membrane microviscosity.

There are many physical and chemical factors that regulate the fluidity properties of biological membranes, including temperature, pressure, membrane potential, fatty acid composition, protein incorporation, and $\text{Ca}^{2+}$ concentration. For example, calcium influenced the structure of membranes containing acidic phospholipids by nonspecifically cross-linking the negative charges. Consequently, increasing the calcium concentration in systems induced structural rearrangements and a decrease in membrane fluidity [12]. Similarly, changes in microfluidity and lateral diffusion fluidity were exhibited when polyunsaturated fatty acids oxidized [13].

Fluidity is an important property of membranes because of its role in various cellular functions. Activities of integral membrane-bound enzymes, such as Na$^+$, K$^+$-ATPase, can be regulated to some extent by changes in the lipid portions of biological membranes. In turn, changes in enzyme activities tightly connected to ion transport processes could affect translocations of ions.

C. Phase Transitions

As is the case for triacylglycerols, phospholipids can exist in a frozen gel state or in a fluid liquid crystalline state, depending on the temperature [14] as illustrated in Fig. 3. Transitions between the gel and liquid crystalline phases can be monitored by a variety of techniques, including nuclear magnetic resonance (NMR), electron spin resonance, fluorescence, and differential scanning calorimetry (DSC). With DSC, both enthalpy and cooperativity of the transition may be determined, enthalpy being the energy required to melt the acyl chains and cooperativity reflecting the number of molecules that undergo a transition simultaneously. However, difficulties in determining membrane transitions have been attributed to entropy/enthalpy com-
pensations in that enthalpy lost by lipids undergoing transition is absorbed by membrane proteins as they partition into the more fluid phase of the bilayer [14].

For complex mixtures of lipids found in biological membranes, at temperatures above the phase transition, all component lipids are liquid crystalline, exhibiting characteristics consistent with complete mixing of the various lipids. At temperatures below the phase transition of the phospholipid with the highest melting temperature, separation of the component into crystalline domains (lateral phase separation) can occur. This ability of individual lipid components to adopt gel or liquid crystalline arrangements has led to the suggestion that particular lipids in a biological membrane may become segregated into a local gel state. This segregation could affect protein function by restricting protein mobility in the bilayer matrix, or it could provide packing defects, resulting in permeability changes. Exposure of plant food tissues to refrigerator temperatures could thus induce localized membrane phase transitions, upset metabolic activity, and create an environment that serves to reduce the quality of the product [15].

Several compositional factors play a role in determining transition temperatures of membranes. The longer the chain length in a phospholipid class, the higher the transition temperature. Similarly, cation binding to PS membranes decreases the phase transition temperature. However, the presence of cis double bonds on the phospholipid fatty acids inhibits hydrocarbon chain packing in the gel state and causes the phase transition to occur at a lower temperature. On the other hand, the presence of the free fatty acid, oleic acid, had negligible effects on the bilayer phase transition, whereas the free fatty acid, palmitic acid, increased the bilayer phase transition temperature [16]. Differential effects on bilayer properties were also seen by the incorporation of cholesterol, and these effects were dependent on the cholesterol concentration [17]. In small amounts (≤3 mol %), a softening of the bilayers in the transition region occurred. However, higher cholesterol concentrations led to a rigidification of the bilayer that was characterized as a liquid-ordered phase. This phase is liquid in the sense that the molecules diffuse laterally as in a fluid, but at the same time the lipid-acyl chains have a high degree of conformational order.

D. Membrane Lipid–Protein Interactions

Complete functioning of a biomembrane is controlled by both the protein and the lipid, mainly phospholipid, components. In a bilayer membrane that contains a heterogeneous distribution of both peripheral and integral proteins, there will be a certain proportion of the phospholipids interacting with the protein component to give the membrane its integrity at both the structural and the functional level. Thus, the proportion of phospholipids in the bilayer interacting with protein at any one time is dictated by protein density, protein type, protein size, and aggregation state of the proteins.

The major structural element of the transmembrane part of many integral proteins is the α-helix bundle and the disposition and packing of such helices determine the degree of protein–lipid interactions. A single α helix passing through a bilayer membrane has a diameter of about 0.8–1 nm, depending on side chain extension, which is similar to the long dimension of the cross-section of a diacyl phospholipid (~0.9–1.0 nm) [18]. In the absence of any significant lateral restriction of such an individual peptide helix, the lateral and rotational motion of the peptide will be
similar to that for the lipids. As the protein mass in or on the membrane increases, however, the motional restriction of the adjacent lipid also increases.

Phospholipids may interact with protein interfaces in selective or nonselective ways. In the absence of selectivity, the lipids act as solvating species, maintaining the protein in a suitable form for activity and mobility. Under these conditions, bilayer fluidity may alter the activity of a membrane protein, with rigid bilayers reducing or inhibiting protein function and fluid bilayers permitting or enhancing protein activity. In selective interactions, associations between polar amino acids and the phospholipid head group may occur with protein function activated by such an interaction.

Phospholipid–protein interactions have important functional consequences. As one example, most ion gradients are set up by active transport proteins, which subsequently are used to drive secondary transport processes. If the ion gradients are lost too quickly by nonspecific leakage (e.g., through membrane regions at the protein–lipid interface), energy will be lost unnecessarily and thus will not be converted to useful work. Another consequence of protein–lipid interactions may be a result of the mutual dynamic influence of one component on the other. It is possible that for biochemical activity to take place, a fluidity window is required within the bilayer part of a membrane for the proteins to undergo the requisite rates and degrees of molecular motion around the active site. When proteins, such as ion-translocating ATPases, undergo significant conformational changes, these rearrangements may not be possible in a solid matrix. Since it is the lipid component of such bilayers that provides this fluidity window, changes in this component can alter the proteins’ activities.

E. Membrane Deterioration and Associated Quality Losses in Food

Quality losses in both plant and animal tissues may be attributed to membrane breakdown following slaughter or harvest. However, postmortem changes in animal tissues occur more rapidly than in plant tissues. In animals, cessation of circulation in the organism leads to lack of oxygen and accumulation of waste products, whereas in plants, respiratory gases can still diffuse across cell membranes and waste products are removed by accumulation in vacuoles.

Two different membrane breakdown pathways predominate in food tissues: free radical lipid oxidation, and loss of plasma and organelle membrane integrity. Some representative modifications that occur in membranes in response to lipid peroxidation include uncoupling of oxidative phosphorylation in mitochondria; alteration of endoplasmic reticulum function; increased permeability; altered activity; inactivation of membrane-bound enzymes, and polymerization, cross-linking, and covalent binding of proteins [19]. Another consequence of lipid peroxidation is formation of the volatile aldehydes that contribute to the aroma characteristics of many vegetables. With regard to loss of plasma and organelle membrane integrity, influx and efflux of solutes may occur, leading to intimate contact among formerly separated catalytic molecules. Thus, in plants where small changes in calcium flux bring about a wide range of physiological responses, catastrophic changes may proceed in the event of loss of membrane integrity. Specific examples of membrane deterioration in both animal and plant tissues are listed in Table 3. A more detailed discussion on these types of membrane deterioration may be found in the review by Stanley [15].
Table 3 Membrane Deterioration in Animal and Plant Tissues

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Description of deterioration</th>
<th>Manifestations of deterioration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Animal</td>
<td>Loss of membrane integrity</td>
<td>Drip</td>
</tr>
<tr>
<td></td>
<td>Oxidative degradation of membrane lipids</td>
<td>Generation of off-flavors: rancid, warmed-over</td>
</tr>
<tr>
<td>Plant</td>
<td>Loss of membrane integrity</td>
<td>Loss of crispness</td>
</tr>
<tr>
<td></td>
<td>Chilling injury</td>
<td>Surface pitting; discoloration</td>
</tr>
<tr>
<td></td>
<td>Senescence/aging</td>
<td>Premature yellowing</td>
</tr>
<tr>
<td></td>
<td>Dehydration</td>
<td>Failure to rehydrate</td>
</tr>
</tbody>
</table>

V. EMULSIFYING PROPERTIES OF PHOSPHOLIPIDS

When one of two immiscible liquid phases is dispersed in the other as droplets, the resulting mixture is referred to as an emulsion. To aid in the stabilization of mainly oil/water emulsions, phospholipids may act as an emulsifier by adsorbing at the interface of the two phases, their amphipathic character contributing to the lowering of interfacial tension. To characterize this process more specifically, a sequence of phase or pseudophase transitions was described near the phase boundary between immiscible liquids upon hydration of an adsorbed phospholipid in \( n \) -decane [20]. These transitions were spherical reverse micelles \( \rightarrow \) three-dimensional network from entangled wormlike micelles \( \rightarrow \) organogel separation into a diluted solution and a compact gel or solid mass precipitating on the interfacial boundary. When prepared in the presence of electrolytes, however, these phospholipid emulsions have a poor stability due to the ability of electrolytes to enhance the vibration of the phospholipid groups at the interface [21]. To circumvent the destabilizing effect of electrolytes, steric surfactants at low concentrations (0.025–0.05%) may be added [22].

Both soybean lecithin and egg yolk are used commercially as emulsifying agents. Egg yolk contains 10% phospholipid and has been used to help form and stabilize emulsions in mayonnaise, salad dressing, and cakes. Commercial soybean lecithin, containing equal amounts of phosphatidylcholine (PC) and inositol, has also been used as an emulsifying agent in ice cream, cakes, candies, and margarine. To expand the range of food grade emulsifiers having different hydrophilic and lipophilic properties, lecithins have been modified physically and enzymatically.

VI. HYDROLYSIS OF PHOSPHOLIPIDS

Several types of ester functionality, all capable of hydrolysis, are present in the component parts of glycerophospholipids (Fig. 1). These may be hydrolyzed totally by chemical methods or selectively by either chemical or enzymatic methods.

A. Chemical Hydrolysis

Mild acid hydrolysis (trichloroacetic acid, acetic acid, HCl, and a little HgCl\(_2\)) results in the complete cleavage of alk-1-enyl bonds of plasmalogens, producing long chain aldehydes. With increasing strength of acid and heating (e.g., 2 N HCl or glacial acetic acid at 100\( ^\circ \)C), diacylglycerol and inositol phosphate are formed from phos-
Phosphatidylinositol (PI) and diacylglycerol and glyceroldiphosphate are formed from diphosphatidyglycerol. Total hydrolysis into each of the component parts of all phospholipids can be accomplished by strong acid (HCl, H₂SO₄) catalysis in 6 N aqueous or 5–10% methanolic solutions [23]. Kinetics and mechanism for hydrolysis in 2 N HCl at 120°C have been described by DeKoning and McMullan [24]. Deacylation occurs first, followed by formation of a cyclic phosphate triester as an intermediate to cyclic glycerophosphate and choline. Eventually an equilibrium mixture of α- and β-glycerophosphates is formed.

Mild alkaline hydrolysis of ester bonds in phospholipids at 37°C (0.025–0.1 M NaOH in methanolic or ethanolic solutions) leads to fatty acids and glycerophosphates. In contrast, phosphosphingolipids are not affected unless subjected to strong alkaline conditions. Some selectivity is seen in the susceptibility of phosphoglycerides to hydrolyze with diacyl > alk-1-enyl, acyl > alkyl, acyl. With more vigorous alkaline hydrolysis, the glycerophosphates are apt to undergo further hydrolysis because the phosphoester bond linking the hydrophilic component to the phospholipid moiety is not stable enough under alkaline conditions and splits, yielding a cyclic phosphate. When the cycle opens up, it gives a 1:1 mixture of 2- and 3-glycerophosphates.

Both state of aggregation and specific polar group have been shown to affect the reaction rates for alkaline hydrolysis of phospholipids [25]. Higher activation energies were observed for hydrolysis of phospholipids in membrane vesicles than when phospholipids were present as monomers or Triton X-100 micelles. Alkaline hydrolysis of PC, on the other hand, was three times faster than hydrolysis of PE.

### B. Enzymatic Hydrolysis

Selective hydrolysis of glycerophospholipids can be achieved by the application of phospholipases. One beneficial aspect to application of phospholipase is improved emulsifying properties to a PC mixture [26]. Unfortunately, while these enzymes may be isolated from a variety of sources, in general they are expensive.

Several phospholipases exist differing in their preferential site of attack. The ester linkage between the glycerol backbone and the phosphoryl group is hydrolyzed by phospholipase C while the ester linkage on the other side of the phosphoryl group is hydrolyzed by phospholipase D. Hydrolysis of the acyl groups at the sn-1 and sn-2 position of phospholipids is carried out by phospholipases A₁ and A₂, respectively.

While phospholipase A₂ binding to membrane phospholipids has been enhanced 10-fold by the presence of calcium [27], membrane surface electrostatics dominated phospholipase A₂ binding and activity in the absence of calcium [28]. A highly cationic enzyme (pI > 10.5), phospholipase A₂, has a marked preference for anionic phospholipid interfaces. Thus, phosphatidic acid and palmitic acid promoted the binding of phospholipase A₂ to the bilayer surface [28,29]. Perturbations and a loosening of the structure associated with the presence of these hydrolysis products were suggested as the properties contributing to enhanced binding [30]. The presence of phospholipid hydroperoxides has also facilitated enhanced binding of phospholipases through a similar mechanism [31].

### VII. HYDROGENATION OF PHOSPHOLIPIDS

Hydrogenation of fats involves the addition of hydrogen to double bonds in the chains of fatty acids. While hydrogenation is more typically applied to triacylglycer-
erols to generate semisolid or plastic fats more suitable for specific applications, it may also be applied to phospholipid fractions. Hydrogenated lecithins are more stable and more easily bleached to a light color, and therefore are more useful as emulsifiers than the natural, highly unsaturated lecithin from soybean oil. These advantages are exemplified by a report that hydrogenated lecithin functions well as an emulsifier and as an inhibitor of fat bloom in chocolate [32].

In practice, hydrogenation involves the mixing of the lipid with a suitable catalyst (usually nickel), heating, and then exposing the mixture to hydrogen at high pressures during agitation. Phospholipids are not as easily hydrogenated as triacylglycerols; as a result, their presence decreases the catalyst activity toward triacylglycerols [33]. In this situation, phosphatidic acid was the most potent poisoning agent; however, fine-grained nickel catalyst was more resistant to the poisoning effect of phospholipids than moderate-grained catalyst. In any event, hydrogenation of phospholipids requires higher temperatures and higher hydrogenation pressures. For example, hydrogenation of lecithin is carried out at 75–80°C in at least 70 atm pressure and in the presence of a flaked nickel catalyst [34]. In chlorinated solvents or in mixtures of these solvents with alcohol, much lower temperatures and pressures can be used for hydrogenation, particularly when a palladium catalyst is used [35].

VIII. HYDROXYLATION

Hydroxylation of the double bonds in the unsaturated fatty acids of lecithin improves the stability of the lecithin and its dispersibility in water and aqueous media. Total hydroxylating agents for lecithin include hydrogen peroxide in glacial acetic acid and sulfuric acid [36]. Such products have been advocated as useful in candy manufacture in which sharp moldings can be obtained when the hydroxylated product is used with starch molds.

IX. HYDRATION

The amount of water absorbed by phospholipids has been measured by a number of different methods, including gravimetry, X-ray diffraction, neutron diffraction, NMR, and DSC [37]. For any measurement, however, Klose et al. [38] cautioned that the morphology and method of sample preparation can induce the formation of defects in and between the bilayers, and therefore will influence the water content of lamellar phospholipids.

The electrical charge on the phospholipid head group does not in itself determine the nature of the water binding. However, it does affect the amount of water bound. The amount of water absorbed by PC from the vapor phase increased monotonically from 0 water molecules per PC molecule at 0% humidity to between 14 [39] and 20 [40] water molecules per lipid molecule at 100% relative humidity. Observed differences may be due to the difficulty of exerting accurate control over relative humidities near 100% when temperature gradients in the system are present. The results of X-ray diffraction studies indicated that when directly mixed with bulk water PC imbibed up to 34 water molecules [41,42]. Considerably less water was imbibed by PE, with a maximum of about 18 water molecules per lipid [43]. From the saturated vapor phase, however, liquid crystalline egg yolk PE only absorbed about 10 water molecules per lipid molecule [39], whereas for charged phospholipids,
such as PI or phosphatidylserine (PS), the phospholipid imbibed water without limit [44,45].

Hydration of a phospholipid appears to be cooperative. A water molecule that initiated hydration of a site facilitated access of additional water molecules, until the hydration of the whole site composed of many different interacting polar residues was completed [46]. Incorporation of the first three to four water molecules on each phospholipid occurs on the phosphate of the lipid head group and is exothermic [47]. The remaining water molecules are incorporated endothermically.

Neutron diffraction experiments on multilayers containing PC [48,49], PE [50], and PI [51] have revealed that water distributions are centered between adjacent bilayers and overlap the head group peaks in the neutron scattering profile of the bilayer. These results imply that water penetrates into the bilayer head group region, but appreciable quantities of water do not reach the hydrocarbon core. By combining X-ray diffraction and dilatometry data, McIntosh and Simon [52,53] were able to calculate the number of water molecules in the interbilayer space and in the head group region for dilauroyl-PE bilayers. They found that there are about 7 and 10 water molecules in the gel and liquid crystalline phases, respectively, with about half of these water molecules located between adjacent bilayers and the other half in the head group region.

The amount of water taken up by a given phospholipid depends on interactions between the lipid molecules, including interbilayer forces (those perpendicular to the plane of the bilayer) and intrabilayer forces (those in the plane of the bilayer). For interbilayer forces, at least four repulsive interactions have been shown to operate between bilayer surfaces. These are the electrostatic, undulation, hydration (solvation), and steric pressures. Attractive pressures include the relatively long-range van der Waals pressure and short-range bonds between the molecules in apposing bilayers, such as hydrogen bonds or bridges formed by divalent salts. Several of the same repulsive and attractive interactions act in the plane of the bilayer, including electrostatic repulsion, hydration repulsion, steric repulsion, and van der Waals attraction. In addition, interfacial tension plays an important role in determining the area per lipid molecule [54]. Thus, as the area per molecule increases, more water can be incorporated into the head group region of the bilayer. Such a situation is found with bilayers having an interdigitated gel phase compared with the normal gel phase and with bilayers having unsaturated fatty acids in the phospholipid compared with saturated fatty acids [55,56].

The presence of monovalent and/or divalent cations in the fluid phase changes the hydration properties of the phospholipids. For example, the partial fluid thickness between dipalmitoyl PC bilayers increased from about 20 Å in water to more than 90 Å in 1 mM CaCl$_2$ [57]. In contrast, monovalent cations, such as Na$^+$, K$^+$, or Cs$^+$, decrease the fluid spaces between adjacent charged PS or PG bilayers as a result of screening of the charge [58,59]. In addition, divalent cations have a dehydrating effect on PS. The most extensively studied divalent cation, Ca$^{2+}$, binds to the phosphate group of PS [60], liberates water between bilayers and from the lipid polar groups [61], crystallizes the lipid hydrocarbon chains [59,60], and raises the gel to the liquid crystalline melting temperature of dipalmitoyl PS by more than 100°C [59]. In response to these changes, one would expect permeability of the membrane to be altered.
The interaction of phospholipids with water is critical to the formation, maintenance, and function of membranes and organelles. It is the low solubility of the acyl chains in water combined with the strong hydrogen bonding between the water molecules that furnishes the “attractive” force that holds together polar lipids as supramolecular complexes (the “hydrophobic bond”). These ordered structures are generated when the phospholipid concentration exceeds its critical micelle concentration (cmc), which is dependent on the free energy gained when an isolated amphiphile in solution enters an aggregate [61]. For diacyl phospholipids in water, the cmc in general is quite low, but it depends on both the chain length and the head group. For a given chain length, the solubility of charged phospholipids is higher, while the cmc of a single-chain phospholipid is higher than that of a diacyl phospholipid with the same head group and the same chain length [61].

X. COMPLEXATION OF PHOSPHOLIPIDS

A. Ions

To comprehend ion binding to phospholipid molecules or to phospholipid membranes, it is necessary to understand the behavior of ions in bulk solution and in the vicinity of a membrane–solution interface. If ion–solvent interactions are stronger than the intermolecular interactions in the solvent, ions are prone to be positively hydrating or structure-making (cosmotropic) entities. The entropy of water is decreased for such ions, while it is increased near other ion types with a low charge density. The latter ions are thus considered to be negatively hydrating or structure-breaking (chaotropic) entities.

When an ion approaches a phospholipid membrane it experiences several forces, the best known of which is the long-range electrostatic, Coulombic force. This force is proportional to the product of all involved charges (on both ions and phospholipids) and inversely proportional to the local dielectric constant. Since phospholipid polar head groups in an aqueous medium are typically hydrated, ion–phospholipid interactions are mediated by dehydration upon binding. Similarly, dehydration of the binding ion may occur. For instance, a strong dehydration effect is observed upon cation binding to the acidic phospholipids, where up to eight water molecules are expelled from the interface once cation–phospholipid association has taken place [62–64].

Various degrees of binding exist between phospholipids and ions. When several water molecules are intercalated between the ion and its binding site, there was actually an association between the ion and phospholipid rather than binding. Outer-sphere complex formation between ion and phospholipid exists when only one water molecule is shared between the ion and its ligand. On the other hand, complete displacement of the water molecules from the region between an ion and its binding site corresponds to an inner-sphere complex. Forces involved in the inner-sphere complex formation include ion–dipole, ion–induced dipole, induced dipole–induced dipole, and ion–quadrupole forces, in addition to the Coulombic interaction. Hydrogen bonding can also participate in inner sphere complex formation. Under appropriate circumstances, the outer-sphere complexes may also be stabilized by “through-water” hydrogen bonding.

Phospholipid affinity for cations appears to follow the sequence lanthanides > transition metals > alkaline earths > alkali metals, thus documenting the significance
of electrostatic interactions in the process of ion–membrane binding. Electrostatic forces also play a strong role in lipid–anion binding with affinity for anions by PC, following the sequence ClO\textsuperscript{−} > I\textsuperscript{−} ≥ SCN\textsuperscript{−} > NO\textsubscript{3}\textsuperscript{−} ≥ Br\textsuperscript{−} > Cl\textsuperscript{−} > SO\textsubscript{4}\textsuperscript{2−}. Here anion size also has an important role in the process of binding, partly as a result of the transfer of the local excess charges from the anion to the phospholipid head groups and vice versa. However, strength of anion binding to phospholipid membranes decreases with increasing net negative charge density of the membrane [58].

Results of NMR, infrared spectroscopy, and neutron diffraction studies strongly imply that the inorganic cations interact predominantly with the phosphodiester groups of the phospholipid head groups [63–67]. On the other hand, inorganic anions may interact specifically with the trimethylammonium residues of the PC head groups [68,69].

Temperature may influence binding of ions to phospholipids. Under conditions of phase transitions, phospholipid chain melting results in a lateral expansion of the lipid bilayers, which for charged systems is also associated with the decrease in the net surface charge density. In the case of negatively charged membranes, this transition leads to lowering of the interfacial proton concentration and decreases the apparent pK value of the anion phospholipids [70].

B. Protein

Complexes of PC with soy protein have been demonstrated by Kanamoto et al. [71]. In this study, using a linear sucrose density gradient centrifugation analysis, \textsuperscript{14}C-PC was found to be nonspecifically bound to either the 7S or 11S proteins.

C. Iodine

In the presence of phospholipid micelles, iodine changes color in aqueous solution. It does not undergo color change in the presence of unassociated molecular species. The color change coincides with the cmc of the substance and is due to formation of the triiodide ion, I\textsubscript{3}−; [72]. Based on data from laser Raman studies, the reaction appeared to be related to iodine–phospholipid interaction, as well as to penetration of iodine into the bilayer membrane, rather than to an ion transport process.

XI. OXIDATION

Unsaturated fatty acids of phospholipids are susceptible to oxidation through both enzymatically controlled processes and random autoxidation processes. The mechanism of autoxidation is basically similar to the oxidative mechanism of fatty acids or esters in the bulk phase or in inert organic solvents. This mechanism is characterized by three main phases: initiation, propagation, and termination. Initiation occurs as hydrogen is abstracted from an unsaturated fatty acid of a phospholipid, resulting in a lipid free radical. The lipid free radical in turn reacts with molecular oxygen to form a lipid peroxyl radical. While irradiation can directly abstract hydrogen from phospholipids, initiation is frequently attributed to reaction of the fatty acids with active oxygen species, such as the hydroxyl free radical and the protonated form of superoxide. These active oxygen species are produced when a metal ion, particularly iron, interacts with triplet oxygen, hydrogen peroxide, and superoxide anion. On the other hand, enzymatic abstraction of hydrogen from an unsaturated
fatty acid occurs when \( \text{Fe}^{3+} \) at the active site of lipoxygenase is reduced to \( \text{Fe}^{2+} \). While the majority of lipoxygenases require free fatty acids, there have been reports of lipoxygenase acting directly on fatty acids in phospholipids [73,74]. Hence, enzymatic hydrolysis may not always be required prior to lipoxygenase activity.

During propagation, lipid–lipid interactions foster propagation of free radicals produced during initiation by abstracting hydrogen from adjacent molecules; the result is a lipid hydroperoxide and a new lipid free radical. Magnification of initiation by a factor of 10 [75] to 100 [76] may occur through free radical chain propagation. Further magnification may occur through branching reactions (also known as secondary initiation) in which \( \text{Fe}^{2+} \) interacts with a hydroperoxide to form a lipid alkoxyl radical and hydroxyl radical, which will then abstract hydrogens from unsaturated fatty acids.

There are many consequences to phospholipid peroxidation in biological and membrane systems. On a molecular level, lipid peroxidation has been manifested in a decreased hydrocarbon core width and molecular volume [77]. In food, the decomposition of hydroperoxides to aldehydes and ketones is responsible for the characteristic flavors and aromas that collectively are often described by the terms “rancid” and “warmed-over.” Numerous studies, on the other hand, have shown that specific oxidation products may be desirable flavor components [78–81], particularly when formed in more precise (i.e., less random) reactions by the action of lipoxygenase enzymes [82–87] and/or by the modifying influence of tocopherol on autoxidation reactions [88].

Through in vitro studies, membrane phospholipids have been shown to oxidize faster than emulsified triacylglycerols [89], apparently because propagation is facilitated by the arrangement of phospholipid fatty acids in the membrane. However, when phospholipids are in an oil state, they are more resistant to oxidation than triacylglycerols or free fatty acids [90]. Evidence that phospholipids are the major contributors to the development of warmed-over flavor in meat from different animal species has been described in several sources [91–94]. Similarly, during frozen storage of salmon fillets, hydrolysis followed by oxidation of the \( \alpha-3 \) fatty acids in phospholipids was noted [95]. The relative importance of phospholipids in these food samples has been attributed to the high degree of polyunsaturation in this lipid fraction and the proximity of the phospholipids to catalytic sites of oxidation (enzymic lipid peroxidation, heme-containing compounds) [96]. However, the importance of phospholipids has not been restricted to animal and fish tissues. In an accelerated storage test of potato granules, both the amounts of phospholipids and their unsaturation decreased [97]. Moreover, with pecans, a much stronger negative correlation was found between headspace hexanal and its precursor fatty acid (18:2) from the phospholipid fraction \((R = -0.98)\) than from the triacylglycerol fraction \((R = -0.66)\) or free fatty acid fraction \((R = -0.79)\) [98]. These results suggest that despite the fact that membrane lipid constitutes a small percentage of the total lipid (0.5%), early stages of oxidation may actually occur primarily within the phospholipids.

The presence of phospholipids does not preclude acceleration of lipid oxidation. When present as a minor component of oil systems, solubilized phospholipids have limited the oxidation of the triacylglycerols [99–101]. Order of effectiveness of individual phospholipids was as follows: \( \text{SPH} = \text{LPC} = \text{PC} = \text{PE} > \text{PS} > \text{PI} > \text{PG} \) [102] with both the amino and hydroxy groups in the side chain participating in the antioxidant activity [103]. It was postulated that antioxidant Maillard reaction prod-
ucts were formed when aldehydes reacted with the amino group of the nitrogen-containing phospholipid. Alternatively, antioxidant activity occurred when complexes between peroxyl free radicals and the amino group were formed [104]. The latter activity is supported by an extended induction period when both tocopherol and phospholipids were present.

Fatty acid composition is a major factor affecting the susceptibility of a phospholipid to assume an oxidized state, with carbon–hydrogen dissociation energies decreasing as the number of bisallylic methylene positions increase [105,106]. However, lipid unsaturation also physically affects oxidation. In model membrane bilayers made from single unilamellar vesicles, lipid unsaturation resulted in smaller vesicles and therefore a larger curvature of the outer bilayer leaflet. The increased lipid–lipid spacing of these highly curved bilayers, in turn, facilitated penetration by oxidants [107,108]. Other functional groups on the phospholipid will also impact their oxidative stability. For example, the presence of an enol ether bond at position 1 of the glycerol backbone in plasmalogen phospholipids has led to inhibition of lipid oxidation, possibly through the binding of the enol ether double bond to initiating peroxyl radicals [109]. Apparently, products of enol ether oxidation do not readily propagate oxidation of polyunsaturated fatty acids. Alternatively, inhibition of lipid oxidation by plasmalogens has been attributed to the iron binding properties of these compounds [110]. Variation within the phospholipid classes toward oxidation has also been ascribed to the iron trapping ability of the polar head group [111]. For example, PS was shown to inhibit lipid peroxidation induced by a ferrous–ascorbate system in the presence of PC hydroperoxides [112]. However, stimulation of phospholipid oxidation by trivalent metal ions (Al$^{3+}$, Sc$^{3+}$, Ga$^{3+}$, In$^{3+}$, Be$^{2+}$, Y$^{3+}$, and La$^{3+}$) has been attributed to the capacity of the ions to increase lipid packing and promote the formation of rigid clusters or displacement to the gel state—processes that bring phospholipid acyl chains closer together to favor propagation steps [113–115].

XII. SUMMARY

This chapter has attempted to highlight the major chemical activities associated with phospholipids and the relevance of these activities to the function of phospholipids in foods. When present in oils or formulated foods, phospholipids may have either detrimental or beneficial effects. As a major component of membranes, phospholipids may also impact the quality of food tissues to a significant extent. Consequently, their modifying presence should not be overlooked, even when they represent a small proportion of the total lipid of a given food tissue.

REFERENCES


Lipid-Based Emulsions and Emulsifiers

D. JULIAN MCCLEMENTS

University of Massachusetts, Amherst, Massachusetts

I. INTRODUCTION

Many natural and processed foods exist either partly or wholly as emulsions, or have been in an emulsified state at some time during their existence [1–5]. Milk is the most common example of a naturally occurring food emulsion [6]. Mayonnaise, salad dressing, cream, ice cream, butter, and margarine are all examples of manufactured food emulsions. Powdered coffee whiteners, sauces, and many desserts are examples of foods that were emulsions at one stage during their production but subsequently were converted into another form. The bulk physicochemical properties of food emulsions, such as appearance, texture, and stability, depend ultimately on the type of molecules the food contains and their interactions with one another. Food emulsions contain a variety of ingredients, including water, lipids, proteins, carbohydrates, minerals, sugars, and small-molecule surfactants [3]. By a combination of covalent and physical interactions, these ingredients form the individual phases and structural components that give the final product its characteristic physicochemical properties [7]. It is the role of food scientists to untangle the complex relationship between the molecular, structural, and bulk properties of foods, so that foods with improved properties can be created in a more systematic fashion.

II. EMULSIONS

An emulsion is a dispersion of droplets of one liquid in another liquid with which it is incompletely miscible [1,8]. In foods, the two immiscible liquids are oil and water. The diameter of the droplets in food emulsions are typically within the range
Emulsions are thermodynamically unstable systems that tend to revert back to the individual oil and water phases with time. To produce an emulsion, energy must be supplied.

A system that consists of oil droplets dispersed in an aqueous phase is called an oil-in-water (O/W) emulsion. A system that consists of water droplets dispersed in an oil phase is called a water-in-oil (W/O) emulsion. The material that makes up the droplets in an emulsion is referred to as the dispersed or internal phase, whereas the material that makes up the surrounding liquid is called the continuous or external phase. Multiple emulsions can be prepared that consist of oil droplets contained in larger water droplets, which are themselves dispersed in an oil phase (O/W/O), or vice versa (W/O/W). Multiple emulsions can be used for protecting certain ingredients, for controlling the release of ingredients, or for creating low-fat products [9].

Emulsions are thermodynamically unstable systems because of the positive free energy required to increase the surface area between the oil and water phases [3]. The origin of this energy is the unfavorable interaction between oil and water, which exists because water molecules are capable of forming strong hydrogen bonds with other water molecules but not with oil molecules [8,9]. Thus emulsions tend to reduce the surface area between the two immiscible liquids by separating into a system that consists of a layer of oil (lower density) on top of a layer of water (higher density). This is clearly seen if one tries to homogenize pure oil and pure water together: initially an emulsion is formed, but after a few minutes phase separation occurs (Fig. 1).

Emulsion instability can manifest itself through a variety of physicochemical mechanisms, including creaming, flocculation, coalescence, and phase inversion (Sec. VI). To form emulsions that are kinetically stable for a reasonable period (a few weeks, months, or even years), chemical substances known as emulsifiers must be added prior to homogenization. Emulsifiers are surface-active molecules that adsorb to the surface of freshly formed droplets during homogenization, forming a protective membrane that prevents the droplets from coming close enough together to aggregate [3]. Most food emulsifiers are amphiphilic molecules, i.e., they have both polar and nonpolar regions on the same molecule. The most common types used in the food industry are lipid-based emulsifiers (small molecule surfactants and phospholipids) and amphiphilic biopolymers (proteins and polysaccharides) [2,3].

Most food emulsions are more complex than the simple three-component (oil, water, and emulsifier) system described above [3,5,9]. The aqueous phase may contain water-soluble ingredients of many different kinds, including sugars, salts, acids, bases, surfactants, proteins, and polysaccharides [1]. The oil phase may contain a variety of lipid-soluble components, such as triacylglycerols, diacylglycerols, monoaoylglycerols, fatty acids, vitamins, and cholesterol [1]. The interfacial membrane

**Figure 1** Emulsions are thermodynamically unstable systems that tend to revert back to the individual oil and water phases with time. To produce an emulsion, energy must be supplied.
may be composed of surface-active components of a variety of types, including small-molecule surfactants, phospholipids, polysaccharides, and proteins. Some of the ingredients in food emulsions are not located exclusively in one phase but are distributed between the oil, water, and interfacial phases according to their partition coefficients. Despite having low concentrations, many of the minor components present in an emulsion can have a pronounced influence on its bulk physicochemical properties. Food emulsions may consist of oil droplets dispersed in an aqueous phase (e.g., mayonnaise, milk, cream, soups), or water droplets dispersed in an oil phase (e.g., margarine, butter, spreads). The droplets and/or the continuous phase may be fluid, gelled, crystalline, or glassy. The size of the droplets may vary from less than a micrometer to a few hundred micrometers, and the droplets themselves may be more or less polydisperse.

To complicate matters further, the properties of food emulsions are constantly changing with time because of the action of various chemical, physical, and biological processes. In addition, during their processing, storage, transport, and handling, food emulsions are subjected to variations in their temperature (e.g., via sterilization, cooking, chilling, freezing) and to various mechanical forces (e.g., stirring, mixing, whipping, flow through pipes, centrifugation high pressure) that alter their physicochemical properties. Despite the compositional, structural, and dynamic complexity of food emulsions, considerable progress has been made in understanding the major factors that determine their bulk physicochemical properties.

III. LIPID-BASED EMULSIFIERS

A. Molecular Characteristics

The most important types of lipid-based emulsifier used in the food industry are small-molecule surfactants (e.g., Tweens, Spans, and salts of fatty acids) and phospholipids (e.g., lecithin). The principal role of lipid-based emulsifiers in food emulsions is to enhance the formation and stability of the product; however, they may also alter the bulk physicochemical properties by interacting with proteins or polysaccharides, or by modifying the structure of fat crystals [9]. All lipid-based emulsifiers are amphiphilic molecules that have a hydrophilic “head” group with a high affinity for water and lipophilic “tail” group with a high affinity for oil [8,10,11]. These emulsifiers can be represented by the formula RX, where X represents the hydrophilic head and R the lipophilic tail. Lipid-based emulsifiers differ with respect to type of head group and tail group. The head group may be anionic, cationic, zwitterionic, or nonionic. The lipid-based emulsifiers used in the food industry are mainly nonionic (e.g., monoacylglycerols, sucrose esters, Tweens, and Spans), anionic (e.g., fatty acids), or zwitterionic (e.g., lecithin). The tail group usually consists of one or more hydrocarbon chains, having between 10 and 20 carbon atoms per chain. The chains may be saturated or unsaturated, linear or branched, aliphatic and/or aromatic. Most lipid-based emulsifiers used in foods have either one or two linear aliphatic chains, which may be saturated or unsaturated. Each type of emulsifier has unique functional properties that depend on its chemical structure.

Lipid-based emulsifiers aggregate spontaneously in solution to form a variety of thermodynamically stable structures known as association colloids (e.g., micelles, bilayers, vesicles, reversed micelles) (Fig. 2). These structural types are adopted
because they minimize the unfavorable contact area between the nonpolar tails of the emulsifier molecules and water [10]. The type of association colloid formed depends principally on the polarity and molecular geometry of the emulsifier molecules (Sec. III.C.3). The forces holding association colloids together are relatively weak, and so they have highly dynamic and flexible structures [8]. Their size and shape is continually fluctuating, and individual emulsifier molecules rapidly exchange between the micelle and the surrounding liquid. The relative weakness of the forces holding association colloids together also means that their structures are particularly sensitive to changes in environmental conditions, such as temperature, pH, ionic strength, and ion type. Surfactant micelles are the most important type of association colloid formed in many food emulsions, and we focus principally on their properties.

**B. Functional Properties**

1. **Critical Micelle Concentration**

   A surfactant forms micelles in an aqueous solution when its concentration exceeds some critical level, known as the critical micelle concentration (cmc). Below the cmc, surfactant molecules are dispersed predominantly as monomers, but once the cmc has been exceeded, any additional surfactant molecules form micelles, and the monomer concentration remains constant. Despite the highly dynamic nature of their structure, surfactant micelles do form particles that have a well-defined average size. Thus, when surfactant is added to a solution above the cmc, the number of micelles increases, rather than their size. When the cmc is exceeded, there is an abrupt change in the physicochemical properties of a surfactant solution (e.g., surface tension, electrical conductivity, turbidity, osmotic pressure) [12]. This is because the properties of surfactant molecules dispersed as monomers are different from those in micelles. For example, surfactant monomers are amphiphilic and have a high surface activity, whereas micelles have little surface activity because their surface is covered with hydrophilic head groups. Consequently, the surface tension of a solution decreases with increasing surfactant concentration below the cmc but remains fairly constant above it.
2. Cloud Point

When a surfactant solution is heated above a certain temperature, known as the cloud point, it becomes turbid. As the temperature is raised, the hydrophilic head groups become increasingly dehydrated, which causes the emulsifier molecules to aggregate. These aggregates are large enough to scatter light, and so the solution appears turbid. At temperatures above the cloud point, the aggregates grow so large that they sediment under the influence of gravity and form a separate phase. The cloud point increases as the hydrophobicity of a surfactant molecule increases; that is, the length of its hydrocarbon tail increases or the size of its hydrophilic head group decreases [13,14].

3. Solubilization

Nonpolar molecules, which are normally insoluble or only sparingly soluble in water, can be solubilized in an aqueous surfactant solution by incorporation into micelles or other types of association colloid [9]. The resulting system is thermodynamically stable; however, equilibrium may take an appreciable time to achieve because of the activation energy associated with transferring a nonpolar molecule from a bulk phase to a micelle. Micelles containing solubilized materials are referred to as swollen micelles or microemulsions, whereas the material solubilized within the micelle is referred to as the solubilizate. The ability of micellar solutions to solubilize nonpolar molecules has a number of potentially important applications in the food industry, including selective extraction of nonpolar molecules from oils, controlled ingredient release, incorporation of nonpolar substances into aqueous solutions, transport of nonpolar molecules across aqueous membranes, and modification of chemical reactions [9]. Three important factors determine the functional properties of swollen micellar solutions: the location of the solubilizate within the micelles, the maximum amount of material that can be solubilized per unit mass of surfactant, and the rate at which solubilization proceeds [9].

4. Surface Activity and Droplet Stabilization

Lipid-based emulsifiers are used widely in the food industry to enhance the formation and stability of food emulsions. To do this they must adsorb to the surface of emulsion droplets during homogenization and form a protective membrane that prevents the droplets from aggregating with each other [1]. Emulsifier molecules adsorb to oil–water interfaces because they can adopt an orientation in which the hydrophilic part of the molecule is located in the water, while the hydrophobic part is located in the oil. This minimizes the unfavorable free energy associated with the contact of hydrophilic and hydrophobic regions, and therefore reduces the interfacial tension. This reduction in interfacial tension is important because it facilitates the further disruption of emulsion droplets; that is, less energy is needed to break up a droplet when the interfacial tension is lowered.

Once adsorbed to the surface of a droplet, the emulsifier must provide a repulsive force that is strong enough to prevent the droplet from aggregating with its neighbors. Ionic surfactants provide stability by causing all the emulsion droplets to have the same electric charge, hence to repel each other electrostatically. Nonionic surfactants provide stability by generating a number of short-range repulsive forces (e.g., steric overlap, hydration, thermal fluctuation interactions) that prevent the drop-
lets from getting too close together [1,11]. Some emulsifiers form multilayers (rather than monolayers) at the surface of an emulsion droplet, which greatly enhances the stability of the droplets against aggregation.

In summary, emulsifiers must have three characteristics to be effective. First, they must rapidly adsorb to the surface of the freshly formed emulsion droplets during homogenization. Second, they must reduce the interfacial tension by a significant amount. Third, they must form a membrane that prevents the droplets from aggregating.

C. Ingredient Selection

A large number of different types of lipid-based emulsifier can be used as food ingredients, and a manufacturer must select the one that is most suitable for each particular product. Suitability, in turn, depends on factors such as an emulsifier’s legal status as a food ingredient; its cost and availability, the consistency in its properties from batch to batch, its ease of handling and dispersion, its shelf life, its compatibility with other ingredients, and the processing, storage, and handling conditions it will experience, as well as the expected shelf life and physicochemical properties of the final product.

How does a food manufacturer decide which emulsifier is most suitable for a product? There have been various attempts to develop classification systems that can be used to select the most appropriate emulsifier for a particular application. Classification schemes have been developed that are based on an emulsifier’s solubility in oil and/or water (Bancroft’s rule), its ratio of hydrophilic to lipophilic groups (HLB number) [15,16], and its molecular geometry [17]. Ultimately, all of these properties depend on the chemical structure of the emulsifier, and so all the different classification schemes are closely related.

1. Bancroft’s Rule

One of the first empirical rules developed to describe the type of emulsion that could be stabilized by a given emulsifier was proposed by Bancroft. Bancroft’s rule states that the phase in which the emulsifier is most soluble will form the continuous phase of an emulsion. Hence, a water-soluble emulsifier will stabilize oil-in-water emulsions, whereas an oil-soluble emulsifier will stabilize water-in-oil emulsions.

2. Hydrophilic–Lipophilic Balance

The hydrophilic–lipophilic balance (HLB) concept underlies a semiempirical method for selecting an appropriate emulsifier or combination of emulsifiers to stabilize an emulsion. The HLB is described by a number, which gives an indication of the overall affinity of an emulsifier for the oil and/or aqueous phases [12]. Each emulsifier is assigned an HLB number according to its chemical structure. A molecule with a high HLB number has a high ratio of hydrophilic groups to lipophilic groups, and vice versa. The HLB number of an emulsifier can be calculated from a knowledge of the number and type of hydrophilic and lipophilic groups it contains, or it can be estimated from experimental measurements of its cloud point. The HLB numbers of many emulsifiers have been tabulated in the literature [15,16]. A widely used semiempirical method of calculating the HLB number of a lipid-based emulsifier is as follows:
HLB = 7 + Σ (hydrophilic group numbers) − Σ (lipophilic group numbers) (1)

As indicated in Table 1 [18], group numbers have been assigned to hydrophilic and lipophilic groups of many types. The sums of the group numbers of all the lipophilic groups and of all the hydrophilic groups are substituted into Eq. (1) and the HLB number is calculated. The semiempirical equation above has been found to have a firm thermodynamic basis, with the sums corresponding to the free energy changes in the hydrophilic and lipophilic parts of the molecule when micelles are formed.

The HLB number of an emulsifier gives a useful indication of its solubility in the oil and/or water phases, and it can be used to predict the type of emulsion that will be formed. An emulsifier with a low HLB number (4–6) is predominantly hydrophobic, dissolves preferentially in oil, stabilizes water-in-oil emulsions, and forms reversed micelles in oil. An emulsifier with a high HLB number (8–18) is predominantly hydrophilic, dissolves preferentially in water, stabilizes oil-in-water emulsions, and forms micelles in water. An emulsifier with an intermediate HLB number (6–8) has no particular preference for either oil or water. Nonionic molecules with HLB numbers below 4 and above 18 are less surface active and are therefore less likely to preferentially accumulate at an oil–water interface.

Emulsion droplets are particularly prone to coalescence when they are stabilized by emulsifiers that have extreme or intermediate HLB numbers. At very high or very low HLB numbers, a nonionic emulsifier has such a low surface activity that it does not accumulate appreciably at the droplet surface and therefore does not provide protection against coalescence. At intermediate HLB numbers (6–8), emulsions are unstable to coalescence because the interfacial tension is so low that very little energy is required to disrupt the membrane. Maximum stability of emulsions is obtained for oil-in-water emulsions using an emulsifier with a HLB number around 10–12, and for water-in-oil emulsions around 3–5. This is because the emulsifiers are sufficiently surface-active but do not lower the interfacial tension so much that the droplets are easily disrupted. It is possible to adjust the effective HLB number by using a combination of two or more emulsifiers with different HLB numbers.

One of the major drawbacks of the HLB concept is its failure to account for the significant alterations in the functional properties of an emulsifier molecule that result from changes in temperature or solution conditions, even though the chemical

<table>
<thead>
<tr>
<th>Hydrophilic group</th>
<th>Group number</th>
<th>Lipophilic group</th>
<th>Group number</th>
</tr>
</thead>
<tbody>
<tr>
<td>—SO₄Na⁺</td>
<td>38.7</td>
<td>—CH—</td>
<td>0.475</td>
</tr>
<tr>
<td>—COOH</td>
<td>21.2</td>
<td>—CH₂—</td>
<td>0.475</td>
</tr>
<tr>
<td>Tertiary amine</td>
<td>9.4</td>
<td>—CH₃</td>
<td>0.475</td>
</tr>
<tr>
<td>Sorbitan ring</td>
<td>6.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>—COOH</td>
<td>2.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>—O—</td>
<td>1.3</td>
<td></td>
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Source: Adopted from Ref. 18.
structure of the molecule does not change. Thus, an emulsifier may be capable of stabilizing oil-in-water emulsions at one temperature but water-in-oil emulsions at another temperature.

3. Molecular Geometry and Phase Inversion Temperature

The molecular geometry of an emulsifier molecule is described by a packing parameter $p$ (see Fig. 3) as follows:

$$p = \frac{v}{l a_o}$$

where $v$ and $l$ are the volume and length of the hydrophobic tail, and $a_o$ is the cross-sectional area of the hydrophilic head group. When surfactant molecules associate with each other, they tend to form monolayers having a curvature that allows the most efficient packing of the molecules. At this optimum curvature, the monolayer has its lowest free energy, and any deviation from this curvature requires the expenditure of energy [8,11]. The optimum curvature of a monolayer depends on the packing parameter of the emulsifier: for $p = 1$, monolayers with zero curvature are preferred; for $p < 1$, the optimum curvature is convex; and for $p > 1$ the optimum curvature is concave (Fig. 3). Simple geometrical considerations indicate that spherical micelles are formed when $p$ is less than 0.33, nonspherical micelles when $p$ is between 0.33 and 0.5, and bilayers when $p$ is between 0.5 and 1 [11]. Above a certain concentration, bilayers join up to form vesicles because energetically unfavorable end effects are eliminated. At values of $p$ greater than 1, reversed micelles are formed, in which the hydrophilic head groups are located in the interior (away from the oil), and the hydrophobic tail groups are located at the exterior (in contact with the oil) (Fig. 2). The packing parameter therefore gives a useful indication of the type of association colloid that is formed by an emulsifier molecule in solution.

The packing parameter is also useful because it accounts for the temperature dependence of the physicochemical properties of surfactant solutions and emulsions. The temperature at which an emulsifier solution converts from a micellar to a reversed micellar system or an oil-in-water emulsion converts to a water-in-oil emul-

![Figure 3](image-url)
sion is known as the phase inversion temperature (PIT). Consider what happens when an emulsion that is stabilized by a lipid-based emulsifier is heated (Fig. 4). At temperatures well below the PIT ($\approx 20^\circ$C), the packing parameter is significantly less than unity, and so a system that consists of oil-in-water emulsion in equilibrium with a swollen micellar solution is favored. As the temperature is raised, the hydrophilic head groups of the emulsifier molecules become increasingly dehydrated, which causes $p$ to increase toward unity. Thus the emulsion droplets become more prone to coalescence and the swollen micelles grow in size. At the phase inversion temperature, $p \approx 1$, and the emulsion breaks down because the droplets have an ultralow interfacial tension and therefore readily coalesce with each other. The resulting system consists of excess oil and excess water (containing some emulsifier monomers), separated by a third phase that contains emulsifier molecules aggregated into bilayer structures. At temperatures sufficiently greater than the PIT, the packing parameter is much larger than unity, and the formation of a system that consists of a water-in-oil emulsion in equilibrium with swollen reversed micelles is favored. A further increase in temperature leads to a decrease in the size of the reversed micelles and in the amount of water solubilized within them. The method of categorizing emulsifier molecules according to their molecular geometry is now widely accepted as the most useful means of determining the types of emulsion they tend to stabilize [17].

4. Other Factors

The classification schemes mentioned above provide information about the type of emulsion an emulsifier tends to stabilize (i.e., O/W or W/O), but they do not provide much insight into the size of the droplets that form during homogenization or the stability of the emulsion droplets once formed [1]. In choosing a suitable emulsifier for a particular application, these factors must also be considered. The speed at which an emulsifier adsorbs to the surface of the emulsion droplets produced during ho-

![Figure 4](image)

**Figure 4** Phase inversion temperature in emulsions.
mogenization determines the minimum droplet size that can be produced: the faster the adsorption rate, the smaller the size. The magnitude and range of the repulsive forces generated by a membrane, and its viscoelasticity, determine the stability of the droplets to aggregation.

IV. BIOPOLYMERS

Proteins and polysaccharides are the two most important biopolymers used as functional ingredients in food emulsions. These biopolymers are used principally for their ability to stabilize emulsions, enhance viscosity, and form gels.

A. Molecular Characteristics

Molecular characteristics of biopolymers, such as molecular weight, conformation, flexibility, and polarity, ultimately determine the properties of biopolymer solutions. These characteristics are determined by the type, number, and sequence of monomers that make up the polymer. Proteins are polymers of amino acids [19], whereas polysaccharides are polymers of monosaccharides [20]. The three-dimensional structures of biopolymers in aqueous solution can be categorized as globular, fibrous, or random coil (Fig. 5). Globular biopolymers have fairly rigid compact structures; fibrous biopolymers have fairly rigid, rodlike structures; and random coil biopolymers have highly dynamic and flexible structures. Biopolymers can also be classified according to the degree of branching of the chain. Most proteins have linear chains, whereas polysaccharides can have either linear (e.g., amylose) or branched (e.g., amylopectin) chains.

The conformation of a biopolymer in solution depends on the relative magnitude of the various types of attractive and repulsive interaction that occur within and between molecules, as well as the configurational entropy of the molecule. Biopolymers that have substantial proportions of nonpolar groups tend to fold into globular structures in which the nonpolar groups are located in the interior (away from the water) and the polar groups are located at the exterior (in contact with the water) because this arrangement minimizes the number of unfavorable contacts between hydrophobic regions and water. However, since stereochemical constraints and the influence of other types of molecular interaction usually make it impossible for all the nonpolar groups to be located in the interior, the surfaces of globular biopolymers have some hydrophobic character. Many kinds of food protein have compact globular structures, including β-lactoglobulin, α-lactalbumin, and bovine serum albumin [6]. Biopolymers that contain a high proportion of polar monomers, distributed fairly evenly along their backbone, often have rodlike conformations with substantial amounts of helical structure stabilized by hydrogen bonding. Such biopolymers (e.g., collagen, cellulose) usually have low water solubilities because they tend to associate.

![Figure 5](https://example.com/figure5.png)

**Figure 5** Typical molecular conformations adopted by biopolymers in aqueous solution.
strongly with each other rather than with water; consequently, they often have poor functional properties. However, if the chains are branched, the molecules may be prevented from getting close enough together to aggregate, and so they may exist in solution as individual molecules. Predominantly polar biopolymers containing monomers that are incompatible with helix formation (e.g., β-casein) tend to form random coil structures.

In practice, biopolymers may have some regions along their backbone that have one type of conformation and others that have a different conformation. Biopolymers may also exist as isolated molecules or as aggregates in solution, depending on the relative magnitude of the biopolymer–biopolymer, biopolymer–solvent, and solvent–solvent interactions. Biopolymers are also capable of undergoing transitions from one type of conformation to another in response to environmental changes such as alterations in their pH, ionic strength, solvent composition, and temperature. Examples include helix ⇔ random coil and globular ⇔ random coil. In many food biopolymers, this type of transition plays an important role in determining the functional properties (e.g., gelation).

B. Functional Properties

1. Emulsification

Biopolymers that have a high proportion of nonpolar groups tend to be surface-active, i.e., they can accumulate at oil–water interfaces [1–4]. The major driving force for adsorption is the hydrophobic effect. When the biopolymer is dispersed in an aqueous phase, some of the nonpolar groups are in contact with water, which is a thermodynamically unfavorable condition. By adsorbing to an interface, the biopolymer can adopt a conformation of nonpolar groups in contact with the oil phase (away from the water) and hydrophilic groups located in the aqueous phase (in contact with the water). In addition, adsorption reduces the number of contacts between the oil and water molecules at the interface, thereby reducing the interfacial tension. The conformation a biopolymer adopts at an oil–water interface, and the physicochemical properties of the membrane formed, depend on its molecular structure. Flexible random coil biopolymers adopt an arrangement in which the predominantly nonpolar segments protrude into the oil phase, the predominantly polar segments protrude into the aqueous phase, and the neutral regions lie flat against the interface (Fig. 6, left). The membranes formed by molecules of these types tend to

Figure 6 The conformation and unfolding of biopolymers at oil–water interfaces depends on their molecular structure.
have relatively open structures, to be relatively thick, and to have low viscoelastic-
ities. Globular biopolymers (usually proteins) adsorb to an interface so that the pre-
dominantly nonpolar regions on their surface face the oil phase; thus they tend to
have a definite orientation at an interface (Fig. 6, right). Once they have adsorbed
to an interface, biopolymers often undergo structural rearrangements that permit them
to maximize the number of contacts between nonpolar groups and oil [4].

Random coil biopolymers have flexible conformations and therefore rearrange
their structures rapidly, whereas globular biopolymers are more rigid and therefore
unfold more slowly. The unfolding of a globular protein at an interface often exposes
amino acids that were originally located in the hydrophobic interior of the molecule,
which can lead to enhanced interactions with neighboring protein molecules through
hydrophobic attraction or disulfide bond formation. Consequently, globular proteins
tend to form relatively thin and compact membranes, high in viscoelasticity. Thus,
membranes formed from globular proteins tend to be more resistant to rupture than
those formed from random coil proteins [3].

To be effective emulsifiers, biopolymers must rapidly adsorb to the surface of
the emulsion droplets formed during homogenization and provide a membrane that
prevents the droplets from aggregating. Biopolymer membranes can stabilize emul-
sion droplets against aggregation by a number of different physical mechanisms [1].
All biopolymers are capable of providing short-range steric repulsive forces that are
usually strong enough to prevent droplets from getting sufficiently close together to
coalesce. If the membrane is sufficiently thick, it can also prevent droplets from
flocculating. Otherwise, it must be electrically charged so that it can prevent floc-
culation by electrostatic repulsion. The properties of emulsions stabilized by charged
biopolymers are particularly sensitive to the pH and ionic strength of aqueous so-
lutions [1a]. At pH values near the isoelectric point of proteins, or at high ionic
strengths, the electrostatic repulsion between droplets may not be large enough to
prevent the droplets from aggregating (see Sec. VI.A.5).

Proteins are commonly used as emulsifiers in foods because many of them
naturally have a high proportion of nonpolar groups. Most polysaccharides are so
hydphilic that they are not surface-active. However, a small number of naturally
occurring polysaccharides have some hydrophobic character (e.g., gum arabic) or
have been chemically modified to introduce nonpolar groups (e.g., some hydrophob-
ically modified starches), and these biopolymers can be used as emulsifiers.

2. Thickening and Stabilization

The second major role of biopolymers in food emulsions is to increase the viscosity
of the aqueous phase [1a]. This modifies the texture and mouthfeel of the food
product (“thickening”), as well as reducing the rate at which particles sediment or
cream (“stabilization”). Both proteins and polysaccharides can be used as thickening
agents, but polysaccharides are usually preferred because they can be used at much
lower concentrations. The biopolymers used to increase the viscosity of aqueous
solutions are usually highly hydrated and extended molecules or molecular aggre-
gates. Their ability to increase the viscosity depends principally on their molecular
weight, degree of branching, conformation, and flexibility. The viscosity of a dilute
solution of particles increases as the concentration of particles increases [3]:
\[ \eta = \eta_0 (1 + 2.5 \phi) \]  

where \( \eta \) is the viscosity of the solution, \( \eta_0 \) is the viscosity of the pure solvent, and \( \phi \) is the volume fraction of particles in solution.

Biopolymers are able to enhance the viscosity of aqueous solutions at low concentrations because they have an effective volume fraction that is much greater than their actual volume fraction [1a]. A biopolymer rapidly rotates in solution because of its thermal energy, and so it sweeps out a spherical volume of water that has a diameter approximately equal to the end-to-end length of the molecule (Fig. 7). The volume of the biopolymer molecule is only a small fraction of the total volume of the sphere swept out, and so the effective volume fraction of a biopolymer is much greater than its actual volume fraction. Consequently, small concentrations of biopolymer can dramatically increase the viscosity of a solution [Eq. (3)]. The effectiveness of a biopolymer at increasing the viscosity increases as the volume fraction it occupies within the sphere it sweeps out decreases. Thus large, highly extended linear biopolymers increase the viscosity more effectively than small compact or branched biopolymers.

In a dilute biopolymer solution the individual molecules (or aggregates) do not interact with each other. When the concentration of biopolymer increases above some critical value \( c^* \), the viscosity increases rapidly because the spheres swept out by the biopolymers overlap with each another. This type of solution is known as a semidilute solution, because even though the molecules are interacting with one another, each individual biopolymer is still largely surrounded by solvent molecules. At still higher polymer concentrations, the molecules pack so close together that they become entangled, and the system has more gel-like characteristics. Biopolymers that are used to thicken the aqueous phase of emulsions are often used in the semidilute concentration range [3].

Solutions containing extended biopolymers often exhibit strong shear-thinning behavior; that is, their apparent viscosity decreases with increasing shear stress. Some

Figure 7  Extended biopolymers in aqueous solutions sweep out a large volume of water as they rotate, which increases their effective volume fraction and therefore their viscosity.
biopolymer solutions even have a characteristic yield stress. When a stress is applied below the yield stress, the solution acts like an elastic solid, but when it exceeds the yield stress the solution acts like a liquid. Shear thinning tends to occur because the biopolymer molecules become aligned with the shear field, or because the weak physical interactions responsible for biopolymer–biopolymer interactions are disrupted. The characteristic rheological behavior of biopolymer solutions plays an important role in determining their functional properties in food emulsions. For example, a salad dressing must be able to flow when it is poured from a container, but must maintain its shape under its own weight after it has been poured onto a salad. The amount and type of biopolymer used must therefore be carefully selected to provide a low viscosity when the salad dressing is poured (high applied stress), but a high viscosity when the salad dressing is allowed to sit under its own weight (low applied stress).

The viscosity of biopolymer solutions is also related to the mouthfeel of a food product. Liquids that do not exhibit extensive shear-thinning behavior at the shear stresses experienced in the mouth are perceived as being “slimy.” On the other hand, a certain amount of viscosity is needed to contribute to the “creaminess” of a product.

The shear-thinning behavior of biopolymer solutions is also important for determining the stability of food emulsions to creaming [1a]. As oil droplets move through an emulsion, they exert very small shear stresses on the surrounding liquid. Consequently, they experience a very large viscosity, which greatly slows down the rate at which they cream and therefore enhances stability. Many biopolymer solutions also exhibit thixotropic behavior (i.e., their viscosity decreases with time when they are sheared at a constant rate) as a result of disruption of the weak physical interactions that cause biopolymer molecules to aggregate. A food manufacturer must therefore select an appropriate biopolymer or combination of biopolymers to produce a final product that has a desirable mouthfeel and texture.

3. Gelation

Biopolymers are used as functional ingredients in many food emulsions (e.g., yogurts, cheeses, desserts, egg and meat products) because of their ability to cause the aqueous phase to gel [1a]. Gel formation imparts desirable textural and sensory attributes, as well as preventing the droplets from creaming. A biopolymer gel consists of a three-dimensional network of aggregated or entangled biopolymers that entraps a large volume of water, giving the whole structure some solid-like characteristics. The appearance, texture, water-holding capacity, reversibility, and gelation temperature of biopolymer gels depends on the type, structure, and interactions of the molecules they contain.

Gels may be transparent or opaque, hard or soft, brittle or rubbery, homogeneous or heterogeneous; they may exhibit syneresis or have good water-holding capacity. Gelation may be induced by a variety of different methods, including altering the temperature, pH, ionic strength, or solvent quality; adding enzymes; and increasing the biopolymer concentration. Biopolymers may be cross-linked by covalent and/or noncovalent bonds.

It is convenient to distinguish between two types of gel: particulate and filamentous (Fig. 8). Particulate gels consist of biopolymer aggregates (particles or clumps) that are assembled together to form a three-dimensional network. This type
Biopolymer molecules or aggregates can form various types of gel structure, such as particulate or filamentous.

of gel tends to be formed when there are strong attractive forces over the whole surface of the individual biopolymer molecules. Particulate gels are optically opaque because the particles scatter light, and they are prone to syneresis because the large interparticle pore sizes means that the water is not held tightly in the gel network by capillary forces. Filamentous gels consist of filaments of individual or aggregated biopolymer molecules that are relatively thin and tend to be formed by biopolymers that can form junction zones only at a limited number of sites on the surface of a molecule, or when the attractive forces between the molecules are so strong that they stick firmly together and do not undergo subsequent rearrangement [3]. Filamentous gels tend to be optically transparent because the filaments are so thin that they do not scatter light significantly, and they tend to have good water-holding capacity because the small pore size of the gel network means that the water molecules are held tightly by capillary forces.

In some foods a gel is formed upon heating (heat-setting gels), while in others it is formed upon cooling (cold-setting gels). Gels may also be either thermoreversible or thermoirreversible, depending on whether gelation is or is not reversible. Gelatin is an example of a cold-setting thermoreversible gel: when a solution of gelatin molecules is cooled below a certain temperature, a gel is formed, but when it is reheated the gel melts. Egg white is an example of a heat-setting thermoirreversible gel: when an egg is heated above a temperature at which gelation occurs, a characteristic white gel is formed; when the egg is cooled back to room temperature, however, the gel remains white (i.e., it does not revert back to its earlier liquid form). Whether a gel is reversible or irreversible depends on the changes in the molecular structure and organization of the molecules during gelation. Biopolymer gels that are stabilized by noncovalent interactions and do not involve large changes in the structure of the individual molecules prior to gelation tend to be reversible. On the other hand, gels that are held together by covalent bonds or involve large changes in the structure of the individual molecules prior to gelation tend to form irreversible gels.

The type of force holding the molecules together in gels varies from biopolymer to biopolymer. Some proteins and polysaccharides (e.g., gelatin, starch) form helical junction zones through extensive hydrogen bond formation. This type of junction zone tends to form when a gel is cooled, becoming disrupted when it is heated, and thus it is responsible for cold-setting gels. Below the gelatin temperature, the attractive hydrogen bonds favor junction zone formation, but above this temperature the configurational entropy favors a random coil type of structure. Biopolymers with
extensive nonpolar groups (e.g., caseins, denatured whey proteins) tend to associate via hydrophobic interactions. Electrostatic interactions play an important role in determining the gelation behavior of many biopolymers, and so gelation is particularly sensitive to the pH and ionic strength of the solution containing the biopolymers. For example, at pH values sufficiently far from their isoelectric point, proteins may be prevented from gelling because of the electrostatic repulsion between the molecules. However, if the pH of the same solution is adjusted near to the isoelectric point, or salt is added, the proteins gel.

The addition of multivalent ions, such as Ca$^{2+}$, can promote gelation of charged biopolymer molecules by forming salt bridges between the molecules. Proteins with thiol groups are capable of forming covalent linkages through thiol–disulfide interchanges, which help to strengthen and enhance the stability of gels. The tendency for a biopolymer to form a gel under certain conditions, and the physical properties of the gel formed, depend on a delicate balance of biopolymer–biopolymer, biopolymer–solvent, and solvent–solvent interactions of various kinds.

C. Ingredient Selection

A wide variety of proteins and polysaccharides are available as ingredients in foods, each with its own unique functional properties and optimum range of applications. Food manufacturers must decide which biopolymer is the most suitable for each type of food product. The selection of the most appropriate ingredient is often the key to success of a particular product. The factors a manufacturer must consider include the desired properties of the final product (appearance, rheology, mouthfeel, stability), the composition of the product, and the processing, storage, and handling conditions the food will experience during its lifetime, as well as the cost, availability, consistency from batch to batch, ease of handling, dispersibility, and functional properties of the biopolymer ingredient.

V. EMULSION FORMATION

The formation of an emulsion may involve a single step or a number of consecutive steps, depending on the nature of the starting material, the desired properties of the end product, and the instrument used to create it [1a]. Before separate oil and aqueous phases are converted to an emulsion, it is usually necessary to disperse the various ingredients into the phase in which they are most soluble. Oil-soluble ingredients, such as certain vitamins, coloring agents, antioxidants, and surfactants, are mixed with the oil, while water-soluble ingredients, such as proteins, polysaccharides, sugars, salts, and some vitamins, coloring agents, antioxidants, and surfactants, are mixed with the water. The intensity and duration of the mixing process depends on the time required to solvate and uniformly distribute the ingredients. Adequate solvation is important for the functionality of a number of food components. If the lipid phase contains any crystalline material, it is usually necessary to warm it before homogenization to a temperature at which all the fat melts; otherwise it is difficult, if not impossible, to efficiently create a stable emulsion.

The process of converting two immiscible liquids to an emulsion is known as homogenization, and a mechanical device designed to carry out this process is called a homogenizer. To distinguish the nature of the starting material, it is convenient to
Figure 9 The homogenization process can be divided into two steps: primary homogenization (creating an emulsion from two separate phases) and secondary homogenization (reducing the size of the droplets in a preexisting emulsion).
scattered from the emulsion droplets). With time, the system rapidly reverts back to its initial state—a layer of oil sitting on top of the water. This is because the droplets formed during the application of the mechanical agitation are constantly moving around and frequently collide and coalesce with neighboring droplets. As this process continues, the large droplets formed rise to the top of the container and merge together to form a separate layer.

To form a stable emulsion, one must prevent the droplets from merging after they have been formed. This is achieved by having a sufficiently high concentration of a surface-active substance, known as an emulsifier, present during the homogenization process. The emulsifier rapidly adsorbs to the droplet surfaces during homogenization, forming a protective membrane that prevents the droplets from coming close enough together to coalesce. One of the major objectives of homogenization is to produce droplets as small as possible because this usually increases the shelf life of the final product. It is therefore important for the food scientist to understand the factors that determine the size of the droplets produced during homogenization.

It should be noted that homogenization is only one step in the formation of a food emulsion, and many of the other unit operations (e.g., pasteurization, cooking, drying, freezing, whipping) also affect the final quality of the product.

A. Physical Principles of Emulsion Formation

The size of the emulsion droplets produced by a homogenizer depends on a balance between two opposing mechanisms: droplet disruption and droplet coalescence (Fig. 10). The tendency for emulsion droplets to break up during homogenization depends on the strength of the interfacial forces that hold the droplets together, compared to the strength of the disruptive forces in the homogenizer. In the absence of any applied external forces, emulsion droplets tend to be spherical because this shape minimizes the contact area between oil and water phases. Changing the shape of a droplet, or breaking it into smaller droplets, increases this contact area and therefore requires the input of energy. The interfacial force holding a droplet together is given by the Laplace pressure ($\Delta P_l$):
\[ \Delta P_i = \frac{2\gamma}{r} \]  

where \( \gamma \) is the interfacial tension between oil and water, and \( r \) is the droplet radius. This equation indicates that it is easier to disrupt large droplets than small ones and that the lower the interfacial tension, the easier it is to disrupt a droplet. The nature of the disruptive forces that act on a droplet during homogenization depends on the flow conditions (i.e., laminar, turbulent, or cavitational) the droplet experiences and therefore on the type of homogenizer used to create the emulsion. To deform and disrupt a droplet during homogenization, it is necessary to generate a stress that is greater than the Laplace pressure and to ensure that this stress is applied to the droplet long enough to enable it to become disrupted [21–23].

Emulsions are highly dynamic systems in which the droplets continuously move around and frequently collide with each other. Droplet–droplet collisions are particularly rapid during homogenization because of the intense mechanical agitation of the emulsion. If droplets are not protected by a sufficiently strong emulsifier membrane, they tend to coalesce during collision. Immediately after the disruption of an emulsion droplet during homogenization, there is insufficient emulsifier present to completely cover the newly formed surface, and therefore the new droplets are more likely to coalesce with their neighbors. To prevent coalescence from occurring, it is necessary to form a sufficiently concentrated emulsifier membrane around a droplet before it has time to collide with its neighbors. The size of droplets produced during homogenization therefore depends on the time taken for the emulsifier to be adsorbed to the surface of the droplets (\( \tau_{\text{adsorption}} \)) compared to the time between droplet–droplet collisions (\( \tau_{\text{collision}} \)). If \( \tau_{\text{adsorption}} \ll \tau_{\text{collision}} \), the droplets are rapidly coated with emulsifier as soon as they are formed and are stable; but if \( \tau_{\text{adsorption}} \gg \tau_{\text{collision}} \), the droplets tend to rapidly coalesce because they are not completely coated with emulsifier before colliding with one of their neighbors. The values of these two times depend on the flow profile the droplets experience during homogenization, as well as the physicochemical properties of the bulk phases and the emulsifier [1a,23].

B. Role of Emulsifiers

The preceding discussion has highlighted two of the most important roles of emulsifiers during the homogenization process:

1. Their ability to decrease the interfacial tension between oil and water phases and thus reduce the amount of energy required to deform and disrupt a droplet [Eq. (4)]. It has been demonstrated experimentally that when the movement of an emulsifier to the surface of a droplet is not rate-limiting (\( \tau_{\text{adsorption}} \ll \tau_{\text{collision}} \)), there is a decrease in the droplet size produced during homogenization with a decrease in the equilibrium interfacial tension [24].

2. Their ability to form a protective membrane that prevents droplets from coalescing with their neighbors during a collision.

The effectiveness of emulsifiers at creating emulsions containing small droplets depends on a number of factors: (a) the concentration of emulsifier present relative to the dispersed phase; (b) the time required for the emulsifier to move from the bulk phase to the droplet surface; (c) the probability that an emulsifier molecule will
be adsorbed to the surface of a droplet during a droplet-emulsifier encounter (i.e., the adsorption efficiency); (d) the amount by which the emulsifier reduces the interfacial tension; and (e) the effectiveness of the emulsifier membrane at protecting the droplets against coalescence.

It is often assumed that small emulsifier molecules adsorb to the surface of emulsion droplets during homogenization more rapidly than larger ones. This assumption is based on the observation that small molecules diffuse to the interface more rapidly than larger ones under quiescent conditions [3]. It has been demonstrated that under turbulent conditions large surface-active molecules tend to accumulate at the droplet surface during homogenization preferentially to smaller ones [23].

C. Homogenization Devices

There are a wide variety of food emulsions, and each one is created from different ingredients and must have different final characteristic properties. Consequently, a number of homogenization devices have been developed for the chemical production of food emulsions, each with its own particular advantages and disadvantages, and each having a range of foods to which it is most suitably applied [1a]. The choice of a particular homogenizer depends on many factors, including the equipment available, the site of the process (i.e., a factory or a laboratory), the physicochemical properties of the starting materials and final product, the volume of material to be homogenized, the throughput, the desired droplet size of the final product, and the cost of purchasing and running the equipment. The most important types of homogenizer used in the food industry are discussed in the subsections that follow.

1. High Speed Blenders

High speed blenders are the most commonly used means of directly homogenizing bulk oil and aqueous phases. The oil and aqueous phase are placed in a suitable container, which may contain as little as a few milliliters or as much as several liters of liquid, and agitated by a stirrer that rotates at high speeds. The rapid rotation of the blade generates intense velocity gradients that cause disruption of the interface between the oil and water, intermingling of the two immiscible liquids, and breakdown of larger droplets to smaller ones [25]. Baffles are often fixed to the inside of the container to increase the efficiency of the blending process by disrupting the flow profile. High speed blenders are particularly useful for preparing emulsions with low or intermediate viscosities. Typically they produce droplets that are between 1 and 10 μm in diameter.

2. Colloid Mills

The separate oil and water phases are usually blended together to form a coarse emulsion premix prior to their introduction into a colloid mill because this increases the efficiency of the homogenization process. The premix is fed into the homogenizer, where it passes between two disks separated by a narrow gap. One of the disks is usually stationary, while the other rotates at a high speed, thus generating intense shear stresses in the premix. These shear stresses are large enough to cause the droplets in the coarse emulsion to be broken down. The efficiency of the homogenization process can be improved by increasing the rotation speed, decreasing the
flow rate, decreasing the size of the gap between the disks, and increasing the surface roughness of the disks. Colloid mills are more suitable than most other types of homogenizer for homogenizing intermediate or high viscosity fluids (e.g., peanut butter, fish or meat pastes), and they typically produce emulsions with droplet diameters between 1 and 5 μm.

3. High Pressure Valve Homogenizers

Like colloid mills, high pressure valve homogenizers are more efficient at reducing the size of the droplets in a coarse emulsion premix than at directly homogenizing two separate phases [26]. The coarse emulsion premix is forced through a narrow orifice under high pressure, which causes the droplets to be broken down because of the intense disruptive stresses (e.g., impact forces, shear forces, cavitation, turbulence) generated inside the homogenizer [27]. Decreasing the size of the orifice increases the pressure the emulsion experiences, which causes a greater degree of droplet disruption and therefore the production of smaller droplets. Nevertheless, the throughput is reduced and more energy must be expended. A food manufacturer must therefore select the most appropriate homogenization conditions for each particular application, depending on the compromise between droplet size, throughput, and energy expenditure. High pressure valve homogenizers can be used to homogenize a wide variety of food products, ranging from low viscosity liquids to viscoelastic pastes, and can produce emulsions with droplet sizes as small as 0.1 μm.

4. Ultrasonic Homogenizers

A fourth type of homogenizer utilizes high intensity ultrasonic waves that generate intense shear and pressure gradients. When applied to a sample containing oil and water, these waves cause the two liquids to intermingle and the large droplets formed to be broken down to smaller ones. There are two types of ultrasonic homogenizer commonly used in the food industry: piezoelectric transducers and liquid jet generators [28]. Piezoelectric transducers are most commonly found in the small benchtop ultrasonic homogenizers used in many laboratories. They are ideal for preparing small volumes of emulsion (a few milliliters to a few hundred milliliters), a property that is often important in fundamental research when expensive components are used. The ultrasonic transducer consists of a piezoelectric crystal contained in some form of protective metal casing, which is tapered at the end. A high intensity electrical wave is applied to the transducer, which causes the piezoelectric crystal inside to oscillate and generate an ultrasonic wave. The ultrasonic wave is directed toward the tip of the transducer, where it radiates into the surrounding liquids, generating intense pressure and shear gradients (mainly due to cavitational affects) that cause the liquids to be broken up into smaller fragments and intermingled with one another. It is usually necessary to irradiate a sample with ultrasound for a few seconds to a few minutes to create a stable emulsion. Continuous application of ultrasound to a sample can cause appreciable heating, and so it is often advantageous to apply the ultrasound in a number of short bursts.

Ultrasonic jet homogenizers are used mainly for industrial applications. A stream of fluid is made to impinge on a sharp-edged blade, which causes the blade to rapidly vibrate, thus generating an intense ultrasonic field that breaks up any droplets in its immediate vicinity though a combination of cavitation, shear, and turbulence [28]. This device has three major advantages: it can be used for contin-
uous production of emulsions; it can generate very small droplets; and it is more energy efficient than high pressure valve homogenizers (since less energy is needed to form droplets of the same size).

5. Microfluidization

Microfluidization is a technique that is capable of creating an emulsion with small droplet sizes directly from the individual oil and aqueous phases [29]. Separate streams of an oil and an aqueous phase are accelerated to a high velocity and then made to simultaneously impinge on a surface, which causes them to be intermingled and leads to effective homogenization. Microfluidizers can be used to produce emulsions that contain droplets as small as 0.1 μm.

6. Membrane Homogenizers

Membrane homogenizers form emulsions by forcing one immiscible liquid into another through a glass membrane that is uniform in pore size. The size of the droplets formed depends on the diameter of the pores in the membrane and on the interfacial tension between the oil and water phases [30]. Membranes can be manufactured with different pore diameters, with the result that emulsions with different droplet sizes can be produced [30]. The membrane technique can be used either as a batch or a continuous process, depending on the design of the homogenizer. Increasing numbers of applications for membrane homogenizers are being identified, and the technique can now be purchased for preparing emulsions in the laboratory or commercially. These instruments can be used to produce oil-in-water, water-in-oil, and multiple emulsions. Membrane homogenizers have the ability to produce emulsions with very narrow droplet size distributions, and they are highly energy efficient, since there is much less energy loss due to viscous dissipation.

7. Energy Efficiency of Homogenization

The efficiency of the homogenization process can be calculated by comparing the energy required to increase the surface area between the oil and water phases with the actual amount of energy required to create an emulsion. The difference in free energy between the two separate immiscible liquids and an emulsion can be estimated by calculating the amount of energy needed to increase the interfacial area between the oil and aqueous phases (ΔG = γΔA). Typically, this is less than 0.1% of the total energy input into the system during the homogenization process because most of the energy supplied to the system is dissipated as heat, owing to frictional losses associated with the movement of molecules past one another [23]. This heat exchange accounts for the significant increase in temperature of emulsions during homogenization.

8. Choosing a Homogenizer

The choice of a homogenizer for a given application depends on a number of factors, including volume of sample to be homogenized, desired throughput, energy requirements, nature of the sample, final droplet size distribution required, equipment available, and initial and running costs. Even after the most suitable homogenization technique has been chosen, the operator must select the optimum processing conditions, such as temperature, time, flow rate, pressure, valve gaps, rotation rates, and sample composition. If an application does not require that the droplets in an emul-
sion be particularly small, it is usually easiest to use a high speed blender. High speed blenders are also used frequently to produce the coarse emulsion premix that is fed into other devices.

To create an emulsion that contains small droplets (<1 μm), either industrially or in the laboratory, it is necessary to use one of the other methods. Colloid mills are the most efficient type of homogenizer for high viscosity fluids, whereas high pressure valve, ultrasonic, or microfluidization homogenizers are more efficient for liquids that are low or intermediate in viscosity. In fundamental studies one often uses small volumes of sample, and therefore a number of laboratory homogenizers have been developed that are either scaled-down versions of industrial equipment or instruments specifically designed for use in the laboratory. For studies involving ingredients that are limited in availability or expensive, an ultrasonic piezoelectric transducer can be used because it requires only small sample volumes. When it is important to have monodisperse emulsions, the use of a membrane homogenizer would be advantageous.

D. Factors That Determine Droplet Size

The food manufacturer is often interested in producing emulsion droplets that are as small as possible, using the minimum amount of energy input and the shortest amount of time. The size of the droplets produced in an emulsion depends on many different factors, some of which are summarized below [27–30].

*Emulsifier concentration.* Up to a certain level, the size of the droplets usually decreases as the emulsifier concentration increases; above this level, droplet size remains constant. When the emulsifier concentration exceeds the critical level, the size of the droplets is governed primarily by the energy input of the homogenization device.

*Emulsifier type.* At the same concentration, different types of emulsifier produce different sized droplets, depending on their surface load, the speed at which they reach the oil–water interface, and the ability of the emulsifier membrane to prevent droplet coalescence.

*Homogenization conditions.* The size of the emulsion droplets usually decreases as the energy input or homogenization time increases.

*Physicochemical properties of bulk liquids.* The homogenization efficiency depends on the physicochemical properties of the lipids that comprise an emulsion (e.g., their viscosity, interfacial tension, density, or physical state).

VI. EMULSION STABILITY

Emulsions are thermodynamically unstable systems that tend, with time, to separate back into individual oil and water phases (Fig. 1). The term “emulsion stability” refers to the ability of an emulsion to resist changes in its properties with time: the greater the emulsion stability, the longer the time taken for the emulsion to alter its properties [1a]. Changes in the properties of emulsions may be the result of physical processes that cause alterations in the spatial distribution of the ingredients (e.g., creaming, flocculation, coalescence, phase inversion) or chemical processes that cause alterations in the chemical structure of the ingredients (e.g., oxidation, hydrolysis). It is important for food scientists to elucidate the relative importance of each
of these mechanisms, the relationship between them, and the factors that affect them, so that effective means of controlling the properties of food emulsions can be established.

A. Droplet–Droplet Interactions

The bulk properties of food emulsions are largely determined by the interaction of the droplets with each other. If the droplets exert a strong mutual attraction, they tend to aggregate, but if they are strongly repelled they tend to remain as separate entities. The overall interaction between droplets depends on the magnitude and range of a number of different types of attractive and repulsive interaction. A knowledge of the origin and nature of these interactions is important because it enables food scientists to predict and control the stability and physicochemical properties of food emulsions.

Droplet–droplet interactions are characterized by an interaction potential $\Delta G(s)$, which describes the variation of the free energy with droplet separation. The overall interaction potential between emulsion droplets is the sum of various attractive and repulsive contributions [3]:

$$\Delta G(s) = \Delta G_{\text{VDW}}(s) + \Delta G_{\text{electrostatic}}(s) + \Delta G_{\text{hydrophobic}}(s) + \Delta G_{\text{short range}}(s)$$

where $\Delta G_{\text{VDW}}$, $\Delta G_{\text{electrostatic}}$, $\Delta G_{\text{hydrophobic}}$, and $\Delta G_{\text{short range}}$ refer to the free energies associated with van der Waals, electrostatic, hydrophobic, and various short-range forces, respectively. In certain systems, there are additional contributions to the overall interaction potential from other types of mechanism, such as depletion or bridging [1a,1b]. The stability of food emulsions to aggregation depends on the shape of the free energy versus separation curve, which is governed by the relative contributions of the different types of interaction [1–3].

1. van der Waals Interactions

The van der Waals interactions act between emulsion droplets of all types and are always attractive. At close separations, the van der Waals interaction potential between two emulsion droplets of equal radius $r$ separated by a distance $s$ is given by the following equation [12]:

$$\Delta G_{\text{VDW}}(s) = -\frac{A r}{12 s}$$

where $A$ is the Hamaker parameter, which depends on the physical properties of the oil and water phases. This equation provides a useful insight into the nature of the van der Waals interaction. The strength of the interaction decreases with the reciprocal of droplet separation, and so van der Waals interactions are fairly long range compared to other types of interaction. In addition, the strength of the interaction increases as the size of the emulsion droplets increases. In practice, Eq. (6) tends to overestimate the attractive forces because it ignores the effects of electrostatic screening, radiation, and the presence of the droplet membrane on the Hamaker parameter [11].
2. Electrostatic Interactions

Electrostatic interactions occur only between emulsion droplets that have electrically charged surfaces (e.g., those established by ionic surfactants or biopolymers). The electrostatic interaction between two droplets at close separation is given by the following relationship [5]:

\[
\Delta G_{\text{electrostatic}}(s) = 4.3 \times 10^{-9} r \psi_0^2 \ln(1 + e^{-4.5})
\]

where 

\[
\kappa^{-1} = \left( \frac{\varepsilon_0 \varepsilon_r k T}{e^2 \sum c_i z_i^2} \right)^{1/2}
\]

Here \(\kappa^{-1}\) is the thickness of the electric double layer, \(c_i\) and \(z_i\) are the molar concentration and valency of ions of species \(i\), \(\varepsilon_0\) is the dielectric constant of a vacuum, \(\varepsilon_r\) is the relative dielectric constant of the medium surrounding the droplet, \(e\) is the electrical charge, \(\psi_0\) is the surface potential, \(k\) is the Boltzmann constant, and \(T\) is the temperature. These equations provide a useful insight into the nature of the electrostatic interactions between emulsion droplets. Usually all the droplets in food emulsions have the same electrical charge, hence repel each other. Electrostatic interactions are therefore important for preventing droplets from aggregating. The strength of the interactions increases as the magnitude of the surface potential increases; thus the greater the number of charges per unit area at a surface, the greater the protection against aggregation. The strength of the repulsive interaction decreases as the concentration of valency of ions in the aqueous phase increases because counterions “screen” the charges between droplets, which causes a decrease in the thickness of the electrical double layer. Emulsions stabilized by proteins are particularly sensitive to the pH and ionic strength of the aqueous solution, since altering pH changes \(\psi_0\) and altering ionic strength changes \(\kappa^{-1}\). The strength of the electrostatic interaction also increases as the size of the emulsion droplets increases.

3. Hydrophobic Interactions

The surfaces of emulsion droplets may not be completely covered by emulsifier molecules, or the droplet membrane may have some nonpolar groups exposed to the aqueous phase [1a]. Consequently, there may be attractive hydrophobic interactions between nonpolar groups and water. The interaction potential energy per unit area between two hydrophobic surfaces separated by water is given by:

\[
\Delta G_{\text{hydrophobic}}(s) = -0.69 \times 10^{-10} r \phi \exp \left( -\frac{s}{\lambda_0} \right)
\]

where \(\phi\) is the fraction of the droplet surface (which is hydrophobic) and the decay length \(\lambda_0\) is of the order of 1–2 nm [11]. The hydrophobic attraction between droplets with nonpolar surfaces is fairly strong and relatively long range [11]. Hydrophobic interactions therefore play an important role in determining the stability of a number of food emulsions. Protein-stabilized emulsions often have nonpolar groups on the protein molecules exposed to the aqueous phase, and therefore hydrophobic interactions are important. They are also important during homogenization because the droplets are not covered by emulsifier molecules.
4. Short-Range Forces

When two emulsion droplets come sufficiently close together, their interfacial layers start to interact. A number of short-range forces result from these interactions, including steric (osmotic and elastic components), hydration, protrusion, and undulation forces [11,12]. Some progress has been made in developing theories to predict the magnitude and range of short-range forces associated with interfacial layers of fairly simple geometry. Nevertheless, both magnitude and range of these forces are particularly sensitive to the size, shape, conformation, packing, interactions, mobility, and hydration of the molecules in the adsorbed layer, and so it is difficult to predict their contribution to the overall interaction potential with any certainty. Even so, they are usually repulsive and tend to increase strongly as the interfacial layers overlap.

5. Overall Interaction Potential

It is often difficult to accurately calculate the contribution of each type of interaction to the overall interdroplet pair potential because information about the relevant physicochemical properties of the system is lacking. Nevertheless, it is informative to examine the characteristics of certain combinations of interactions that are particularly important in food emulsions, for this provides a valuable insight into the factors that affect the tendency of droplets to aggregate. Consider an emulsion in which the only important types of droplet–droplet interaction are van der Waals attraction, electrostatic repulsion, and steric repulsion (e.g., an emulsion stabilized by a charged biopolymer).

The van der Waals interaction potential is fairly long range and always negative (attractive), the electrostatic interaction potential is fairly long range and always positive (repulsive), while the steric interaction is short range and highly positive (strongly repulsive). The overall interdroplet pair potential has a complex dependence on separation because it is the sum of these three different interactions, and it may be attractive at some separations and repulsive at others. Figure 11 shows a typical

![Figure 11](image)

**Figure 11** The overall interaction potential for an emulsion stabilized by a charged biopolymer.
profile of interdroplet pair potential versus separation for an emulsion stabilized by a charged biopolymer. When the two droplets are separated by a large distance, there is no effective interaction between them. As they move closer together, the van der Waals attraction dominates initially and there is a shallow minimum in the profile, which is referred to as the secondary minimum. If the depth of this minimum is large compared to the thermal energy \( \Delta G(s_{\text{min}}^2) \gg kT \), the droplets tend to be flocculated. However, if it is small compared to the thermal energy, the droplets tend to remain unaggregated. At closer separations the repulsive electrostatic interactions dominate, and there is an energy barrier \( \Delta G(s_{\text{max}}) \) that must be overcome before the droplets can come any closer. If this energy barrier is sufficiently large compared to the thermal energy \( \Delta G(s_{\text{max}}) \gg kT \), it will prevent the droplets from falling into the deep primary minimum at close separations. On the other hand, if it is not large compared to the thermal energy, the droplets will tend to fall into the primary minimum, leading to strong flocculation of the droplets. In this situation the droplets would be prevented from coalescing because of the domination of the strong steric repulsion at close separations.

Emulsions that are stabilized by repulsive electrostatic interactions are particularly sensitive to the ionic strength and pH of the aqueous phase \([1a,1b]\). At low ion concentrations there may be a sufficiently high energy barrier to prevent the droplets from getting close enough together to aggregate into the primary minimum. As the ion concentration is increased, the screening of the electrostatic interactions becomes more effective, which reduces the height of the energy barrier. Above a certain ion concentration, the energy barrier is not high enough to prevent the droplets from falling into the primary minimum, and so the droplets become strongly flocculated. This phenomenon accounts for the tendency for droplets to flocculate when salt is added to emulsions stabilized by ionic emulsifiers. The surface charge density of protein-stabilized emulsions decreases as the pH tends toward the isoelectric point, which reduces the magnitude of the repulsive electrostatic interactions between the droplets and also leads to droplet flocculation.

### B. Mechanisms of Emulsion Instability

As mentioned earlier, emulsions are thermodynamically unstable systems that tend with time to revert back to the separate oil and water phases of which they were made. The rate at which this process occurs, and the route that is taken, depend on the physicochemical properties of the emulsion and the prevailing environmental conditions. The most important mechanisms of physical instability are creaming, flocculation, coalescence, Ostwald ripening, and phase inversion. In practice, all these mechanisms act in concert and can influence one another. However, one mechanism often dominates the others, facilitating the identification of the most effective method of controlling emulsion stability.

The length of time an emulsion must remain stable depends on the nature of the food product. Some food emulsions (e.g., cake batters, ice cream mix, margarine premix) are formed as intermediate steps during a manufacturing processes and need remain stable for only a few seconds, minutes, or hours. Other emulsions (e.g., mayonnaise, creme liqueurs) must persist in a stable state for days, months, or even years prior to sale and consumption. Some food processing operations (e.g., the production of butter, margarine, whipped cream, and ice cream) rely on controlled
destabilization of an emulsion. We now turn to a discussion of the origin of the major destabilization mechanisms, the factors that influence them, and methods of controlling them. This type of information is useful for food scientists because it facilitates the selection of the most appropriate ingredients and processing conditions required to produce a food emulsion with particular properties.

1. Creaming and Sedimentation

The droplets in an emulsion have a density different from that of the liquid that surrounds them, and so a net gravitational force acts on them \[1a,1b\]. If the droplets have lower density than the surrounding liquid, they tend to move up, that is, to “cream.” Conversely, if they have a higher density they tend to move down, resulting in what is referred to as sedimentation. Most liquid oils have densities lower than that of water, and so there is a tendency for oil to accumulate at the top of an emulsion and water at the bottom. Thus droplets in an oil-in-water emulsion tend to cream, whereas those in a water-in-oil emulsion tend to sediment. The creaming rate of a single isolated spherical droplet in a viscous liquid is given by the Stokes equation:

\[
\nu = -\frac{2gr^2(\rho_2 - \rho_1)}{9\eta_1}
\]

where \(\nu\) is the creaming rate, \(g\) the acceleration due to gravity, \(\rho\) the density, \(\eta\) the shear viscosity, and the subscripts 1 and 2 refer to the continuous phase and droplet, respectively. The sign of \(\nu\) determines whether the droplet moves up (+) or down (−).

Equation (9) can be used to estimate the stability of an emulsion to creaming. For example, an oil droplet (\(\rho_2 = 910 \text{ kg/m}^3\)) with a radius of 1 \(\mu\text{m}\) suspended in water (\(\eta_1 = 1 \text{ mPa}\cdot\text{s}, \rho_1 = 1000 \text{ kg/m}^3\)) will cream at a rate of about 5 mm/day. Thus one would not expect an emulsion containing droplets of this size to have a particularly long shelf life. As a useful rule of thumb, an emulsion in which the creaming rate is less than about 1 mm/day can be considered to be stable toward creaming [3].

In the initial stages of creaming (Fig. 12), the droplets move upward and a droplet-depleted layer is observed at the bottom of the container. When the droplets reach the top of the emulsion, they cannot move up any further and so they pack together to form the “creamed layer.” The thickness of the final creamed layer depends on the packing of the droplets in it. Droplets may pack very tightly together, or they may pack loosely, depending on their polydispersity and the magnitude of the forces between them. Close-packed droplets will tend to form a thin creamed layer, whereas loosely packed droplets form a thick creamed layer. The same factors that affect the packing of the droplets in a creamed layer determine the nature of the flocs formed (see Sec. VI.B.2). If the attractive forces between the droplets are fairly weak, the creamed emulsion can be redispersed by lightly agitating the system. On the other hand, if an emulsion is centrifuged, or if the droplets in a creamed layer are allowed to remain in contact for extended periods, significant coalescence of the droplets may occur, with the result that the emulsion droplets can no longer be redispersed by mild agitation.

Creaming of emulsion droplets is usually an undesirable process, which food manufacturers try to avoid. Equation (9) indicates that creaming can be retarded by
minimizing the density difference \((\rho_2 - \rho_1)\) between the droplets and the surrounding liquid, reducing the droplet size, or increasing the viscosity of the continuous phase. The Stokes equation is strictly applicable only to isolated rigid spheres suspended in an infinite viscous liquid. Since these assumptions are not valid for food emulsions, the equation must be modified to take into account hydrodynamic interactions, droplet fluidity, droplet aggregation, non-Newtonian aqueous phases, droplet crystallization, the adsorbed layer, and Brownian motion [1a,2].

2. Flocculation and Coalescence

The droplets in emulsions are in continual motion because of their thermal energy, gravitational forces, or applied mechanical forces, and as they move about they collide with their neighbors. After a collision, emulsion droplets may either move apart or remain aggregated, depending on the relative magnitude of the attractive and repulsive forces between them. If the net force acting between the droplets is strongly attractive, they will aggregate, but if it is strongly repulsive they will remain unaggregated. Two types of aggregation are commonly observed in emulsions: flocculation and coalescence. In flocculations (Fig. 12), two or more droplets come together to form an aggregate in which the emulsion droplets retain their individual integrity. Coalescence is the process whereby two or more droplets merge together to form a single larger droplet (Fig. 12). Improvements in the quality of emulsion-based food products largely depend on an understanding of the factors that cause droplets to aggregate. The rate at which droplet aggregation occurs in an emulsion depends on two factors: collision frequency and collision efficiency [1a,1b].

The collision frequency is the number of encounters between droplets per unit time per unit volume. Any factor that increases the collision frequency is likely to increase the aggregation rate. The frequency of collisions between droplets depends on whether the emulsion is subjected to mechanical agitation. For dilute emulsions containing identical spherical particles, the collision frequency \(N\) has been calculated for both quiescent and stirred systems [3]:

![Figure 12 Mechanisms of emulsion instability.](image-url)
\[ N = \frac{4kTn_0^3}{3\eta} \tag{10} \]
\[ N = \frac{16}{3} Gr^n n_0^2 \tag{11} \]

where \( n_0 \) is the initial number of particles per unit volume and \( G \) is the shear rate. The collision efficiency, \( E \), is the fraction of encounters between droplets that lead to aggregation. Its value ranges from 0 (no flocculation) to 1 (fast flocculation) and depends on the interaction potential. The equations for the collision frequency must therefore be modified to take into account droplet–droplet interactions:

\[ N = \frac{4kTn_0^3}{3\eta} E \tag{12} \]

where

\[
E = \int_{2r}^{\infty} \left\{ \exp \left( \frac{\Delta G(x)}{kT} \right) x^{-2} dx \right\}^{-1}
\]

with \( x \) the distance between the centers of the droplets (\( x = 2r + s \)) and \( \Delta G(x) \) the droplet–droplet interaction potential (Sec. VI.A). Emulsion droplets may remain unaggregated, or they may aggregate into the primary or secondary minima depending on \( \Delta G(x) \).

The equations above are applicable only to the initial stages of aggregation in dilute emulsions containing identical spherical particles. In practice, most food emulsions are fairly concentrated systems, and interactions between flocs as well as between individual droplets are important. The equations above must therefore be modified to take into account the interactions and properties of flocculated droplets.

The nature of the droplet–droplet interaction potential also determines the structure of the flocs formed, and the rheology and stability of the resulting emulsion [1a]. When the attractive force between them is relatively strong, two droplets tend to become “locked” together as soon as they encounter each other. This leads to the formation of flocs that have quite open structures [3]. When the attractive forces are not particularly strong, the droplets may “roll around” each other after a collision, which allows them to pack more efficiently to form denser flocs. These two extremes of floc structure are similar to those formed by filamentous and particulate gels, respectively (Fig. 8).

The structure of the flocs formed in an emulsion has a pronounced influence on its bulk physicochemical properties. An emulsion containing flocculated droplets has a higher viscosity than one containing unflocculated droplets, since the water trapped between the flocculated droplets increases the effective diameter (and therefore volume fraction) of the particles (Eq. 3). Flocculated particles also exhibit strong shear thinning behavior: as the shear rate is increased, the viscosity of the emulsion decreases because the flocs are disrupted and so their effective volume fraction decreases. If flocculation is extensive, a three-dimensional network of aggregated particles extends throughout the system and the emulsion has a yield stress that must be overcome before the system will flow. The creaming rate of droplets is also strongly dependent on flocculation. At low droplet concentrations, flocculation increases the creaming rate because the effective size of the particles is increased.
Partial coalescence occurs when two partly crystalline emulsion droplets collide and aggregate because a crystal in one droplet penetrates the other droplet. [Eq. (9)], but at high droplet concentrations, it retards creaming because the droplets are trapped within the three-dimensional network of aggregated emulsion droplets.

In coalescence (Fig. 12), two or more liquid droplets collide and merge into a single larger droplet. Extensive coalescence eventually leads to oiling off, i.e., formation of free oil on the top of an emulsion. Because coalescence involves a decrease in the surface area of oil exposed to the continuous phase, it is one of the principal mechanisms by which an emulsion reverts to its most thermodynamically stable state (Fig. 1). Coalescence occurs rapidly between droplets that are not protected by emulsifier molecules; for example, if one homogenizes oil and water in the absence of an emulsifier, the droplets readily coalesce. When droplets are stabilized by an emulsifier membrane, the tendency for coalescence to occur is governed by the droplet–droplet interaction potential and the stability of the film to rupture. If there is a strong repulsive force between the droplets at close separations, or if the film is highly resistant to rupture, the droplets will tend not to coalesce. Most food emulsions are stable to coalescence, but they become unstable when subjected to high shear forces that cause the droplets to frequently collide with each other or when the droplets remain in contact with each other for extended periods (e.g., droplets in flocs, creamed layers, or highly concentrated emulsions).

3. Partial Coalescence

Normal coalescence involves the aggregation of two or more liquid droplets to form a single larger spherical droplet, but partial coalescence occurs when two or more partially crystalline droplets encounter each other and form a single irregularly shaped aggregate (Fig. 13). The aggregate is irregular in shape because some of the structure of the fat crystal network contained in the original droplets is maintained within it. It has been proposed that partial coalescence occurs when two partially crystalline droplets collide and a crystal from one of them penetrates the intervening membranes and protrudes into the liquid region of the other droplet [1a]. Normally, the crystal would stick out into the aqueous phase, thus becoming surrounded by water; however, when it penetrates another droplet, it is surrounded by oil, and because this arrangement is energetically favorable the droplets remain aggregated. With time the droplets slowly fuse more closely together, with the result that the total surface area of oil exposed to the aqueous phase is reduced. Partial coalescence occurs only when the droplets have a certain ratio of solid fat and liquid oil. If the solid fat content of the droplets is either too low or too high, the droplets will tend not to undergo partial coalescence [5].

![Figure 13](image)

**Figure 13** Partial coalescence occurs when two partly crystalline emulsion droplets collide and aggregate because a crystal in one droplet penetrates the other droplet.
Partial coalescence is particularly important in dairy products because milk fat globules are partially crystalline at temperatures commonly found in foods. The application of shear forces or temperature cycling to cream containing partly crystalline milk fat globules can cause extensive aggregation of the droplets, leading to a marked increase in viscosity (“thickening”) and subsequent phase separation [9]. Partial coalescence is an essential process in the production of ice cream, whipped toppings, butter, and margarine. Oil-in-water emulsions are cooled to a temperature at which the droplets are partly crystalline, and a shear force is then applied that causes droplet aggregation via partial coalescence. In butter and margarine, aggregation results in phase inversion, whereas in ice cream and whipped cream the aggregated fat droplets form a network that surrounds air cells and provides the mechanical strength needed to produce good stability and texture.

4. Ostwald Ripening

Ostwald ripening is the growth of large droplets at the expense of smaller ones [1a]. This process occurs because the solubility of the material in a spherical droplet increases as the size of the droplet decreases:

\[ S(r) = S(\infty) \exp \left( \frac{2\gamma V_m}{RT} \right) \]  

Here \( V_m \) is the molar volume of the solute, \( \gamma \) is the interfacial tension, \( R \) is the gas constant, \( S(\infty) \) is the solubility of the solute in the continuous phase for a droplet with infinite curvature (i.e., a planar interface), and \( S(r) \) is the solubility of the solute when contained in a spherical droplet of radius \( r \). The greater solubility of the material in smaller droplets means that there is a higher concentration of solubilized material around a small droplet than around a larger one. Consequently, solubilized molecules move from small droplets to large droplets because of this concentration gradient, which causes the larger droplets to grow at the expense of the smaller ones. Once steady state conditions have been achieved, the growth in droplet radius with time due to Ostwald ripening is given by

\[ \frac{d\langle r \rangle^3}{dt} = \frac{8\gamma V_m S(\infty) D}{9RT} \]  

where \( D \) is the diffusion coefficient of the material through the continuous phase. This equation assumes that the emulsion is dilute and that the rate-limiting step is the diffusion of the solute molecules across the continuous phase. In practice, most food emulsions are concentrated systems, and so the effects of the neighboring droplets on the growth rate have to be considered. Some droplets are surrounded by interfacial membranes that retard the diffusion of solute molecules in and out of droplets, and in such cases the equation must be modified accordingly. Ostwald ripening is negligible in many foods because triacylglycerols have extremely low water solubilities, and therefore the mass transport rate is insignificant [Eq. (14)]. Nevertheless, in emulsions that contain more water-soluble lipids, such as flavor oils, Ostwald ripening may be important.

5. Phase Inversion

In phase inversion (Fig. 12), a system changes from an oil-in-water emulsion to a water-in-oil emulsion or vice versa. This process usually occurs as a result of some
alteration in the system’s composition or environmental conditions, such as dispersed phase volume fraction, emulsifier type, emulsifier concentration, temperature, or application of mechanical forces. Phase inversion is believed to occur by means of a complex mechanism that involves a combination of the processes that occur during flocculation, coalescence, and emulsion formation. At the point where phase inversion occurs, the system may briefly contain regions of oil-in-water emulsion, water-in-oil emulsion, multiple emulsions, and bicontinuous phases, before converting to its final state.

6. Chemical and Biochemical Stability

Chemical and biochemical reactions of various types (e.g., oxidation, reduction, or hydrolysis of lipids, polysaccharides, and proteins) can cause detrimental changes in the quality of food emulsions. Many of these reactions are catalyzed by specific enzymes that may be present in the food. The reactions that are important in a given food emulsion depend on the concentration, type, and distribution of ingredients, and the thermal and shear history of the food. Chemical and biochemical reactions can alter the stability, texture, flavor, odor, color, and toxicity of food emulsions. Thus it is important to identify the most critical reactions that occur in each type of food so that they can be controlled in a systematic fashion.

VII. CHARACTERIZATION OF EMULSION PROPERTIES

Ultimately, food manufacturers want to produce a high quality product at the lowest possible cost. To achieve this goal they must have a good appreciation of the factors that determine the properties of the final product. This knowledge, in turn, is used to formulate and manufacture a product with the desired characteristics (e.g., appearance, texture, mouthfeel, taste, shelf life). These bulk physicochemical and sensory properties are determined by such molecular and colloidal properties of emulsions as dispersed volume fraction, droplet size distribution, droplet–droplet interactions, and interfacial properties. Consequently, a wide variety of experimental techniques have been developed to characterize the molecular, colloidal, microscopic, and macroscopic properties of food emulsions [1a]. Analytical techniques are needed to characterize the properties of food emulsions in the laboratory, where they are used to improve our understanding of the factors that determine emulsion properties, and in the factory, where they are used to monitor the properties of foods during processing to ensure that the manufacturing process is operating in an appropriate manner. The subsections that follow highlight some of the most important properties of food emulsions and outline experimental techniques for their measurement.

A. Dispersed Phase Volume Fraction

The dispersed phase volume fraction or \( \phi \) is the volume of emulsion droplets \( V_d \) divided by the total volume of the emulsion \( V_e \): \( \phi = V_d/V_e \). The dispersed phase volume fraction determines the relative proportion of oil and water in a product, as well as influencing many of the bulk physicochemical and sensory properties of emulsions, such as appearance, rheology, taste, and stability. For example, an emulsion tends to become more turbid and to have a higher viscosity when the concentration of droplets is increased [1a]. Methods for measuring the dispersed phase
volume fraction of emulsions are outlined in Table 2. Traditional proximate analysis
techniques, such as solvent extraction to determine oil content and oven drying to
determine moisture content, can be used to analyze the dispersed phase volume
fraction of emulsions. Nevertheless, proximate analysis techniques are often destruc-
tive and quite time-consuming to carry out, and are therefore unsuitable for rapid
quality control or on-line measurements. If the densities of the separate oil and
aqueous phases are known, the dispersed phase volume fraction of an emulsion can
simply be determined from a measurement of its density:

$$\phi = \left(\frac{\rho_{\text{emulsion}}}{\rho_{\text{droplet}}}ight)\left(\frac{\rho_{\text{continuous phase}}}{\rho_{\text{continuous phase}}}ight)$$  (15)

The electrical conductivity of an emulsion decreases as the concentration of oil
within it increases, and so instruments based on electrical conductivity can also be
used to determine $\phi$. Light scattering techniques can be used to measure the dispersed
phase volume fraction of dilute emulsions ($\phi < 0.001$), whereas NMR and ultrasound
spectroscopy can be used to rapidly and nondestructively determine $\phi$ of concen-
trated and optically opaque emulsions. A number of these experimental techniques
(e.g., ultrasound, NMR, electrical conductivity, density measurements) are particu-
larly suitable for on-line determination of the composition of food emulsions during
processing.

B. Droplet Size Distribution

The size of the droplets in an emulsion influences many of their sensory and bulk
physicochemical properties, including rheology, appearance, mouthfeel, and stability
[3,5]. It is therefore important for food manufacturers to carefully control the size
of the droplets in a food product and to have analytical techniques to measure droplet
size. Typically, the droplets in a food emulsion are somewhere in the size range of
0.1–50 $\mu$m in diameter.
Food emulsions always contain droplets that have a range of sizes, and so it is usually important to characterize both the average size and the size distribution of the droplets. The droplet size distribution is usually represented by a plot of droplet frequency (number or volume) versus droplet size (radius or diameter). Some of the most important experimental techniques for measuring droplet size distributions are included in Table 2.*

Light-scattering and electrical conductivity techniques are capable of providing a full particle size distribution of a sample in a few minutes. Since, however, these techniques usually require that the droplet concentration be very low ($d < 0.001$), samples must be diluted considerably before analysis. Optical and electron microscopy techniques, which provide the most direct measurement of droplet size distribution, are often time-consuming and laborious to operate, and sample preparation can cause considerable artifacts in the results. In contrast, recently developed techniques based on NMR and ultrasonic spectroscopy can be used to rapidly and non-destructively measure the droplet size distribution of concentrated and optically opaque emulsions [1a]. These techniques are particularly useful for on-line characterization of emulsion properties.

C. Microstructure

The structural organization and interactions of the droplets in an emulsion often play an important role in determining the properties of a food. For example, two emulsions may have the same droplet concentration and size distribution, but very different properties, because of differences in the degree of droplet flocculation. Various forms of microscopy are available for providing information about the microstructure of food emulsions. The unaided human eye can resolve objects that are farther apart than about 0.1 mm (100 $\mu$m). Most of the structural components in food emulsions (e.g., emulsion droplets, surfactant micelles, fat crystals, ice crystals, small air cells, protein aggregates) are much smaller than this lower limit and cannot therefore be observed directly by the eye.

Optical microscopy can be used to study components of size between about 0.5 and 100 $\mu$m. The characteristics of specific components can be highlighted by selectively staining certain ingredients or by using special lenses. Electron microscopy can be used to study components that have sizes down to about 0.5 nm. Atomic force microscopy can be used to provide information about the arrangements and interactions of single atoms or molecules. All these techniques are burdened by sample preparation steps that often are laborious and time-consuming, and subject to alter the properties of the material being examined. Nevertheless, when carried out correctly the advanced microscopic techniques provide extremely valuable information about the arrangement and interactions of emulsion droplets with each other and with the other structural entities found in food emulsions.

D. Physical State

The physical state of the components in a food emulsion often has a pronounced influence on its overall properties [1a]. For example, oil-in-water emulsions are par-

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*A comprehensive review of analytical methods for measuring particle size in emulsions has recently been published [31].
particularly prone to partial coalescence when the droplets contain a certain percentage of crystalline fat (Sec. VI.B). Partial coalescence leads to extensive droplet aggregation, which decreases the stability of emulsions to creaming and greatly increases their viscosity. In water-in-oil emulsions, such as margarine or butter, the formation of a network of aggregated fat crystals provides the characteristic rheological properties. The most important data for food scientists are the temperature at which melting or crystallization begins, the temperature range over which the phase transition occurs, and the value of the solid fat content at any particular temperature. Phase transitions can be monitored by measuring changes in any property (e.g., density, compressibility, heat capacity, absorption or scattering of radiation) that is altered upon conversion of an ingredient from a solid to a liquid (Table 2). The density of a component often changes when it undergoes a phase transition, and so melting or crystallization can be monitored by measuring changes in the density of a sample with temperature or time.

Phase transitions can also be monitored by measuring the amount of heat absorbed or released when a solid melts or a liquid crystallizes, respectively. This type of measurement can be carried out by means of differential thermal analysis or differential scanning calorimetry. These techniques also provide valuable information about the polymorphic form of the fat crystals in an emulsion. More recently, rapid instrumental methods based on NMR and ultrasound have been developed to measure solid fat contents [1a]. These instruments are capable of nondestructively determining the solid fat content of a sample in a few seconds and are extremely valuable analytical tools for rapid quality control and on-line procedures. Phase transitions can be observed in a more direct manner by means of polarized optical microscopy.

E. Creaming and Sedimentation Profiles

Over the past decade, a number of instruments have been developed to quantify the creaming or sedimentation of the droplets in emulsions. Basically the same light scattering, NMR, and ultrasound techniques used to measure the dispersed phase volume fraction or droplet size distributions of emulsions are applied to creaming or sedimentation, but the measurements are carried out as a function of sample height to permit the acquisition of a profile of droplet concentrations or sizes. Techniques based on the scattering of light can be used to study creaming and sedimentation in fairly dilute emulsions. A light beam is passed through a sample at a number of different heights, and the reflection and transmission coefficients are measured and related to the droplet concentration and size. By measuring the ultrasonic velocity or attenuation as a function of sample height and time, it is possible to quantify the rate and extent of creaming in concentrated and optically opaque emulsions. This technique can be fully automated and has the two additional advantages: creaming can be detected before it is visible to the eye, and a detailed creaming profile can be determined rather than a single boundary. By measuring the ultrasound properties as a function of frequency, it is possible to determine both the concentration and size of the droplets as a function of sample height. Thus a detailed analysis of creaming and sedimentation in complex food systems can be monitored noninvasively. Recently developed NMR imaging techniques can also measure the concentration and size of droplets in any region in an emulsion [9]. These ultrasound and NMR techniques will prove particularly useful for understanding the kinetics of
creaming and sedimentation in emulsions and for predicting the long-term stability of food emulsions.

**F. Emulsion Rheology**

The rheology of an emulsion is one of its most important overall physical attributes because it largely determines the mouthfeel, flowability, and stability of emulsions [3]. A variety of experimental techniques are available for measuring the rheological properties of food emulsions. The rheology of emulsions that have low viscosities and act like ideal liquids can be characterized by capillary viscometers. For nonideal liquids or viscoelastic emulsions, more sophisticated instrumental techniques called dynamic shear rheometers are available to measure the relationship between the stress applied to an emulsion and the resulting strain, or vice versa. As well as providing valuable information about the bulk physicochemical properties of emulsions (e.g., texture, flow through pipes), rheological measurements can provide information about droplet–droplet interactions and the properties of any flocs formed in an emulsion.

**G. Interfacial Properties**

Despite comprising only a small fraction of the total volume of an emulsion, the interfacial region that separates the oil from the aqueous phase plays a major role in determining stability, rheology, chemical reactivity, flavor release, and other overall physicochemical properties of emulsions. The most important properties of the interface are the concentration of emulsifier molecules present (the surface load), the packing of the emulsifier molecules, and the thickness, viscoelasticity, electrical charge, and (interfacial) tension of the interface.

A variety of experimental techniques are available for characterizing the properties of oil–water interfaces (Table 2). The surface load is determined by measuring the amount of emulsifier that adsorbs per unit area of oil–water interface. The thickness of an interfacial membrane can be determined by light scattering, neutron scattering, neutron reflection, surface force, and ellipsometry techniques. The rheological properties of the interfacial membrane can be determined by means of the two-dimensional analog of normal rheological techniques. The electrical charge of the droplets in an emulsion determines their susceptibility to aggregation. Experimental techniques based on electrokinetic and electroacoustic techniques are available for determining the charge on emulsion droplets. The dynamic or equilibrium interfacial tension of an oil–water interface can be determined by means of a number of interfacial tension meters, including the Wilhelmy plate, Du Nouy ring, maximum bubble pressure, and pendant drop methods.

**REFERENCES**

29. E. Dickinson and G. Stainsby. Emulsion stability. In: *Advances in Food Emulsions and

I. CHEMISTRY OF WAXES

A. Introduction

The term *waxes* commonly refers to the mixtures of long chain apolar compounds found on the surface of plants and animals. By a strict chemical definition, a wax is the ester of a long chain acid and a long chain alcohol. However, this academic definition is much too narrow both for the wax chemist and for the requirements of industry. The following description from the German Society for Fat Technology [1] better fits the reality:

Wax is the collective term for a series of natural or synthetically produced substances that normally possess the following properties: kneadable at $20^\circ$C, brittle to solid, coarse to finely crystalline, translucent to opaque, relatively low viscosity even slightly above the melting point, not tending to stinginess, consistency and solubility depending on the temperature and capable of being polished by slight pressure.

The collective properties of wax as just defined clearly distinguish waxes from other articles of commerce. Chemically, waxes constitute a large array of different chemical classes, including hydrocarbons, wax esters, sterol esters, ketones, aldehydes, alcohols, and sterols. The chain length of these compounds may vary from $C_2$, as in the acetate of a long chain ester, to $C_{62}$ in the case of some hydrocarbons [2,3].

Waxes can be classified according to their origins as naturally occurring or synthetic. The naturally occurring waxes can be subclassified into animal, vegetable, and mineral waxes. Beeswax, spermaceti, wool grease, and lanolin are important animal waxes. Beeswax, wool grease, and lanolin are by-products of other industries. The vegetable waxes include carnauba wax, the so-called queen of waxes, ouricouri (another palm wax), and candelilla. These three waxes account for the major pro-
portion of the consumption of vegetable waxes. The mineral waxes are further classified into the petroleum waxes, ozokerite, and montan. Based on their chemical structure, waxes represent a very broad spectrum of chemical types from polyethylene, polymers of ethylene oxide, derivatives of montan wax, alkyl esters of monocarboxylic acids, alkyl esters of hydroxy acids, polyhydric alcohol esters of hydroxy acids, Fisher–Tropsch waxes, and hydrogenated waxes, to long chain amide waxes.

We begin with an overview of the diverse class of lipids known as waxes. The discussion presented that follows, which touches on source, structure, function, and biosynthesis, is intended to serve as an entry to the literature, enabling the reader to pursue this topic in greater detail.

B. Properties and Characteristics of Waxes

The ancient Egyptians used beeswax to make writing tablets and models, and waxes are now described as man’s first plastic. Indeed, the plastic property of waxes and cold-flow yield values allow manual working at room temperature, corresponding to the practices of the Egyptians. The melting points of waxes usually vary within the range 40–120°C.

Waxes dissolve in fat solvents, and their solubility is dependent on temperature. They can also wet and disperse pigments, and they can be emulsified with water, which makes them useful in the furniture, pharmaceutical, and food industries. Their combustibility, associated with a low ash content, is important in candle manufacture and solid fuel preparation. Waxes also find application in industry as lubricants and insulators, where their properties as natural plastics, their high flash points, and their high dielectric constants are advantageous.

The physical and technical properties of waxes depend more on molecular structure than on molecular size and chemical constitution. The chemical components of waxes range from hydrocarbons, esters, ketones, aldehydes, and alcohols to acids, mostly as aliphatic long chain molecules. The hydrocarbons in petroleum waxes are mainly alkanes, though some unsaturated and branched chain compounds are found. The common esters are those of saturated acids with 12–28 carbon atoms combining with saturated alcohols of similar chain length. Primary alcohols, acids, and esters have been characterized and have been found to contain an even straight chain of carbon atoms. By contrast, most ketones, secondary alcohols, and hydrocarbons have odd numbers of carbon atoms. The chemical constitution of waxes varies in great degree depending on the origin of the material. A high proportion of cholesterol and lanosterol is found in wool wax. Commercial waxes are characterized by a number of properties. These properties are used in wax grading [4].

1. Physical Properties of Waxes

Color and odor are determined by comparison with standard samples in a molten state. In the National Petroleum Association scale, the palest color is rated 0, while amber colors are rated 8. Refined waxes are usually free from taste, this property being especially important in products such as candelilla when it is used in chewing gum. Melting and softening points are important physical properties. The melting points can be determined by the capillary tube method or the drop point method. The softening point of a wax is the temperature at which the solid wax begins to soften. The penetration property measures the depth to which a needle with a definite top load penetrates the wax sample.
Shrinkage and flash point are two frequently measured physical properties of waxes. The flash point is the temperature at which a flash will occur if a small flame is passed over the surface of the sample. In the liquid state, a molten wax shrinks uniformly until the temperature approaches the solidification point. This property is measured as the percentage shrinkage of the volume.

2. Chemical Properties of Waxes

a. Acid Value. The acid value is the number of milligrams of potassium hydroxide required to neutralize a gram of the wax. It is determined by the titration of the wax solution in ethanol–toluene with 0.5 M potassium hydroxide. Phenolphthalein is normally used as the titration indicator.

\[
\text{Acid value} = \frac{V_w \times 56.104}{w}
\]

where \( V_w \) is the number of milliliters (mL) of potassium hydroxide used in the titration and \( w \) is the mass of wax.

b. Saponification Number. The saponification number is the number of milligrams of potassium hydroxide required to hydrolyze 1 g of wax:

\[
\text{Saponification number} = \frac{(V_b - V_w) \times 56.105}{w}
\]

where \( w \) is the weight of wax sample(s), \( V_b \) the volume (mL) of hydrochloric acid used in the blank, and \( V_w \) the volume (mL) of hydrochloric acid used in the actual analysis. The wax (2 g) is dissolved in hot toluene (910 mL). Alcoholic potassium hydroxide (25 mL of 0.5 M KOH) is added, and the solution is refluxed for 2 hours. A few drops of phenolphthalein are added and the residual potassium hydroxide is titrated with 0.5 M hydrochloric acid. A blank titration is also performed with 25 mL of 0.5 M alcoholic potassium hydroxide plus toluene.

c. Ester Value. Ester value, the difference between the saponification number and the acid value, shows the amount of potassium hydroxide consumed in the saponification of the esters.

d. Iodine Number. The iodine number expresses the amount of iodine that is absorbed by the wax. It is a measure of the degree of unsaturation.

e. Acetyl Number. The acetyl number indicates the milligrams of potassium hydroxide required for the saponification of the acetyl group assimilated in 1 g of wax on acetylation. The difference of this number and the ester value reflects the amount of free hydroxy groups (or alcohol composition) in a wax. The wax sample is first acetylated by acetic anhydride. A certain amount of acetylated wax (about 2 g) is taken out to be saponified with the standard procedure in the measurement of the saponification number. The acetyl number is the saponification number of the acetylated wax.

3. Properties of Important Naturally Occurring Waxes

a. Beeswax. Beeswax is a hard amorphous solid, usually light yellow to amber depending on the source and manufacturing process. It has a high solubility in warm benzene, toluene, chloroform, and other polar organic solvents. Typically, beeswax
has an acid value of 17–36, a saponification number of 90–147, melting point of 60–67°C, an ester number of 64–84, a specific gravity of 0.927–0.970, and an iodine number of 7–16. Pure beeswax consists of about 70–80% of long chain esters, 12–15% of free acids, 10–15% of hydrocarbon, and small amounts of diols and cholesterol esters. Beeswax is one of the most useful and valuable of waxes. Its consumption is not limited to the candle industry, the oldest field of wax consumption. It is also used in electrical insulation and in the food, paper, and rubber industries.

b. Wool Grease and Lanolin. Wool grease is a by-product of the wool industry, and the finest wool grease yields lanolin. Pharmaceutical grade lanolin accounts for about 80% of all wool grease consumption. Wool grease has a melting point of 35–42°C, an acid value of 7–15, a saponification value of 100–110, an ester value of 85–100, a specific gravity of 0.932–0.945, and an iodine value of 22–30.

c. Carnauba Wax. Carnauba wax, “queen of waxes,” is a vegetable wax produced in Brazil. Carnauba wax is hard, amorphous, and tough, with a pleasant smell. It is usually used in cosmetics and by the food industry, in paper coatings, and in making inks. In the food industry, it is a minor component in glazes for candies, gums, and fruit coatings. Carnauba wax is soluble in most polar organic solvents. It contains esters (84–85%), free acids (3–3.5%), resins (4–6%), alcohols (2–3%), and hydrocarbons (1.5–3.0%). Typically, carnauba has an acid value of 2.9–9.7, an ester value of 39–55, a saponification value of 79–95, an iodine value of 7–14, and a melting range of 78–85°C.

d. Candelilla Wax. Candelilla wax is a vegetable wax produced mainly in Mexico. It is used chiefly in the manufacturing of chewing gum and cosmetics, which represent about 40% of the market. It is also used in furniture polish, in the production of lubricants, and in paper coating. Candelilla wax has a specific gravity of 0.98, an acid value of 12–22, a saponification value of 43–65, a melting point of 66–71°C, an ester value of 65–75, and an iodine value of 12–22. The chemical composition of candelilla wax is 28–29% esters, 50–51% hydrocarbon, 7–9% free acids, and small amounts of alcohols and cholesterol.

e. Ozocerite. Ozocerite is a mineral wax found in Galicia, Russia, Iran, and the United States. Most ozocerite consists of hydrocarbons, but the chemical composition varies with the source. Typically ozocerite has an ester value of 56–66, an acid value of 31–38, a saponification value of 87–104, a melting point of 93–89°C, and an iodine value of 14–18. Ozocerite is graded as unbleached (black), single bleached (yellow), and double bleached (white). It is mainly used in making lubricants, lipsticks, polishes, and adhesives.

C. Isolation, Separation, and Analysis of Natural Waxes

Knowledge of the chemical analysis of natural waxes is essential for understanding wax biosynthesis, manufacture, and application. While the chemical compositions of synthetic waxes are constant and depend on the manufacturing process, the natural waxes are much more complicated in chemical composition. In general, natural waxes are isolated by chemical extraction, separated by chromatographic methods, and analyzed by means of mass spectrometry (MS); both gas chromatography (GC) and high performance liquid chromatography (HPLC) techniques are used. The fol-
lowing discussion on chemical analysis is based on an understanding of the general principles of chemical extraction, chromatography, and mass spectrometry. There are numerous textbooks detailing these principles [5–7].

1. Isolation

Natural waxes are mixtures of long chain apolar compounds found on the surface of plants and animals. However internal lipids also exist in most organisms. In earlier times, the plant or animal tissue was dried, whereupon the total lipid material could be extracted with hexane or chloroform by means of a Soxhlet extractor. The time of exposure to the organic solvent, particularly chloroform, is kept short to minimize or avoid the extraction of internal lipids. Because processors are interested in the surface waxes, it became routine to harvest them by a dipping procedure. For plants this was usually done in the cold, but occasionally at the boiling point of light petroleum or by swabbing to remove surface lipids. Chloroform, which has been widely used, is now known to be toxic; dichloromethane can be substituted. After removal of the solvent under vacuum, the residue can be weighed. Alternatively, the efficiency of the extraction can be determined by adding a known quantity of a standard wax component (not present naturally in the sample) and performing a quantification based on this component following column chromatography.

2. Separation

The extract of surface lipids contains hydrocarbons, as well as long chain alcohols, aldehydes and ketones, short chain acid esters of the long chain alcohols, fatty acids, sterols and sterol esters, and oxygenated forms of these compounds. In most cases it is necessary to separate the lipid extract into lipid classes prior to the identification of components. Separation of waxes into their component classes is first achieved by column chromatography. The extract residue is redissolved in the least polar solvent possible, usually hexane or light petroleum, and transferred to the chromatographic column. When the residue is not soluble in hexane or light petroleum, a hot solution or a more polar solvent, like chloroform or dichloromethane, may be used to load the column. By gradually increasing the polarity of the eluting solvent, it is possible to obtain hydrocarbons, esters, aldehydes and ketones, triglycerides, alcohols, hydroxydiacetones, sterols, and fatty acids separately from the column. Most separations have been achieved on alumina or silica gel. However, Sephadex LH-20 was used to separate the alkanes from Green River Shale. Linde 5-Å sieve can remove the \( n \)-alkanes to provide concentrated branched and alicyclic hydrocarbons. Additionally, silver nitrate can be impregnated into alumina or silica gel columns or thin-layer chromatography (TLC) plates for separating components according to the degree of unsaturation.

As the means of further identifying lipids become more sophisticated, it is possible to obtain a sufficient quantity of the separated wax components by TLC. One of the major advantages of TLC is that it can be modified very easily, and minor changes to the system have allowed major changes in separation to be achieved. Most components of wax esters can be partially or completely separated by TLC on 25 \( \mu \text{m} \) silica gel G plates developed in hexane–diethyl ether or benzene–hexane. The retardation factor \( (R_f) \) values of most wax components are listed in Table 1 [8].

If TLC is used, the components must be visualized, and the methods employed can be either destructive or nondestructive. The commonly used destructive method
Table 1  TLC Separation of Wax Components on Silica Gel: \( R_f \) Values for Common Wax Components

<table>
<thead>
<tr>
<th>Component</th>
<th>Solvent systems&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
</tr>
<tr>
<td>Hydrocarbon</td>
<td>0.95</td>
</tr>
<tr>
<td>Squalene</td>
<td></td>
</tr>
<tr>
<td>Trialkylglyceryl ethers</td>
<td>0.90</td>
</tr>
<tr>
<td>Steryl esters</td>
<td>0.90</td>
</tr>
<tr>
<td>Wax esters</td>
<td>0.90</td>
</tr>
<tr>
<td>( \beta )-Diketones</td>
<td></td>
</tr>
<tr>
<td>Monoketones</td>
<td></td>
</tr>
<tr>
<td>Fatty acid methyl esters</td>
<td>0.65</td>
</tr>
<tr>
<td>Aldehydes</td>
<td>0.55</td>
</tr>
<tr>
<td>Triterpenyl acetates</td>
<td></td>
</tr>
<tr>
<td>Secondary alcohols</td>
<td></td>
</tr>
<tr>
<td>Triacylglycerols</td>
<td>0.35</td>
</tr>
<tr>
<td>Free fatty acids</td>
<td>0.18</td>
</tr>
<tr>
<td>Triterpenols</td>
<td></td>
</tr>
<tr>
<td>Primary alcohols</td>
<td>0.15</td>
</tr>
<tr>
<td>Sterols</td>
<td>0.10</td>
</tr>
<tr>
<td>Hydroxy-( \beta )-diketones</td>
<td>0.09</td>
</tr>
<tr>
<td>Triterpenoid acid</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> A, petroleum ether (b.p. 60–70°C)–diethyl ether–glacial acetic acid (90:10:1, v/v); B, benzene; C, chloroform containing 1% ethanol; D, petroleum ether (b.p. 40–60°C)–diethyl ether (80:20, v/v); E, chloroform containing 1% ethanol; F, hexane–heptane–diethyl ether–glacial acetic acid (63:18.5:18.5, v/v) to 2 cm from top, then full development with carbon tetrachloride; G (1) petroleum ether–diethyl ether–glacial acetic acid (80:20:1, v/v); (2) petroleum ether; H, benzene–chloroform (70:30 v/v).

is to spray TLC plates with sulfuric or molybdic acid in ethanol and heat them. This technique is very sensitive, but it destroys the compounds and does not work well with free acids. Iodine vapors will cause a colored band to appear, particularly with unsaturated compounds, and is widely used to both locate and quantify the lipids. Since the iodine can evaporate from the plate readily after removal from iodine chamber, the components usually remain unchanged. Iodine vapor is one of the ideal visualization media in the isolation of lipid classes from TLC plates. Commercial TLC plates with fluorescent indicators are available as well, and bands can be visualized under UV light. However, if it is necessary to use solvents more polar than diethyl ether to extract polar components from the matrix, the fluorescent indicators may also be extracted, and these additives will interfere with subsequent analyses.

To isolate lipid classes from TLC plates after a nondestructive method of visualization, the silica gel can be scraped into a champagne funnel and eluted with an appropriate solvent. Or, the gel can be scraped into a test tube and the apolar lipid extracted with diethyl ether by vortexing, centrifuging, and decanting off the ether. Polar lipids are extracted in the same manner, using a more polar solvent such as chloroform and/or methanol. High performance liquid chromatography has been used in the separation and analysis of natural waxes, but its application was halted by the lack of a suitable detector, since most wax components have no useful ultra-
violet chromophore. Application of ultraviolet detection was limited to wavelength around 210 nm. Some components with isolated double bonds and carbonyl group (e.g., esters, aldehydes, ketones) can be detected in this wavelength. Hamilton and coworkers have examined an alternative detection system, infrared detection at 5.74 μm, which allowed the hydrocarbon components to be detected [9]. While the sensitivity of this method of detection could not match that of ultraviolet detection, it has merit for use in the preparative mode, where it is feasible to allow the whole output from the column to flow through the detector. The third useful mode for HPLC is mass spectrometry. The coupling of HPLC and MS makes this form of chromatography a very important analytical technique.

3. Analysis

When individual classes of waxes have been isolated, the identity of each must be determined. Because of the complex composition of these materials, combined analytical approaches (e.g., GC-MS) have been used to analyze individual wax classes. Mass spectrometry is a major analytical method for the analysis of this class of compounds. With the electron impact–mass spectrometry (EI-MS), the wax molecules tend to give cleavage fragments rather than parent ions. Thus, soft (chemical) ionization (CI), and fast atom bombardment (FAB) have been frequently used to give additional information for wax analysis.

In GC-MS analysis, the hydrocarbon fraction and many components of the wax ester fraction can be analyzed directly, while long chain alcohols, the aldehydes, and fatty acids are often analyzed as their acetate esters of alcohols, dimethylhydrazones of aldehydes, and methyl ethers of fatty acids. The analysis of wax esters after hydrolysis and derivatization will provide additional information on high molecular weight esters. For example, the chain branching of a certain component might be primarily examined with respect to its unusual retention time on GC analysis, then determined by converting to the corresponding hydrocarbon through the reduction of its iodide intermediate with LiAID₄ (the functional group end is labeled by the deuterium atom). A similar approach is to convert the alcohol of the target component to an alkyl chloride via methanesulfonyl chloride. This method labels the functional end with a chlorine atom, and its mass spectra are easily interpreted because of the chlorine isotopes. As mentioned above, unsaturated hydrocarbons can be separated from saturated hydrocarbons and unsaturated isomers by column chromatography or TLC with silver nitrate silica gel or alumina gel media. The position and number of double bonds affect the volatility of the hydrocarbons, thereby altering their retention in GC and HPLC analysis. The location of a double bond is based on the mass spectra of their derivatives, using either positive or negative CI.

D. Biosynthesis of Natural Waxes

Epicuticular waxes (from the outermost layer of plant and insect cuticles) comprise very long chain nonpolar lipid molecules that are soluble in organic solvents. In many cases this lipid layer may contain proteins and pigments, and great variability in molecular architecture is possible, depending on the chemical composition of the wax and on environmental factors [10,11].

A variety of waxes can be found in the cuticle. On the outer surface of plants these intracuticular waxes entrap cutin, which is an insoluble lipid polymer of hy-
droxy and epoxy fatty acids. In underlying layers, associated with the suberin matrix, another cutin-like lipid polymer containing aliphatic and aromatic components is found [12]. In some instances, internal nonsubarin waxes, which are stored in plant seeds, are the major energy reserves rather than triacylglycerols. In insects, intracuticular waxes are the major constituents of the inner epicuticular layer [13–15].

A variety of aliphatic lipid classes occur in epicuticular waxes. These include hydrocarbons, alcohols, esters, ketones, aldehydes, and free fatty acids of numerous types [16,17]. Frequently, a series of 10 carbon atom homologs occurs, while chain lengths of 10–35 carbon atoms are most often found. However, fatty acids and hydrocarbons with fewer than 20 carbon atoms are known, as are esters with more than 60 carbon atoms. Other minor lipids such as terpenoids, flavonoids, and sterols also occur in epicuticular waxes. The composition and quantity of epicuticular wax varies widely from one species to another and from one organ, tissue, or cell type to another [16]. In insects, wax composition depends on stage of life cycle, age, sex, and external environment [17].

In waxes, the biosynthesis of long chain carbon skeletons is accomplished by a basic condensation–elongation mechanism. Elongases are enzyme complexes that repetitively condense short activated carbon chains to an activated precursor and prepare the growing chain for the next addition. The coordinated action of two such soluble complexes is plastid results in the synthesis of the 16 and 18 carbon acyl chains characterizing plant membranes [18–20]. Each condensation introduces a β-keto group into the elongating chain. This keto group is normally removed by a series of three reactions: a β-keto reduction, a β-hydroxy dehydration, and an enol reduction.

Variations of the foregoing basic biosynthetic mechanism occur, giving rise to compounds classified as polyketides. Their modified acyl chains can be recognized by the presence of keto groups, hydroxy groups, or double bonds that were not removed before the next condensation took place. It is well established that the very long carbon skeletons of the wax lipids are synthesized by a condensation–elongation mechanism. The primary elongated products in the form of free fatty acids are often minor components of epicuticular waxes. Most of them, however, serve as substrates for the associated enzyme systems discussed. The total length attained during elongation is reflected by the chain lengths of the members of the various wax classes [15–21]. Normal, branched, and unsaturated hydrocarbons and fatty acids are prominent components of plant waxes, while insect waxes usually lack long chain free fatty acids [22–26].

II. CHEMISTRY OF STEROLS

A. Introduction

Sterols constitute a large group of compounds with a broad range of biological activities and physical properties. The natural occurring sterols usually possess the 1,2-cyclopentano-phenanthrene skeleton with a stereochemistry similar to the trans-syn-trans-anti-trans-anti configuration at their ring junctions, and have 27–30 carbon atoms with an hydroxy group at C-3 and a side chain of at least seven carbons at C-17 (Fig. 1). Sterols can exhibit both nuclear variations (differences within the ring system) and side chain variations. The examples of the three subclasses of sterols in Figure 1 represent the major variations of sterols. Sterols have been defined as hy-
droxylated steroids that retain some or all of the carbon atoms of squalene in the side chain and partition almost completely into an ether layer when shaken with equal volumes of water and ether [27].

Sterols are common in eukaryotic cells but rare in prokaryotes. Without exception, vertebrates confine their sterol biosynthetic activity to producing cholesterol. Most invertebrates do not have the enzymatic machinery for sterol biosynthesis and must rely on an outside supply. Sterols of invertebrates have been found to comprise most complex mixtures arising through food chains. In plants, cholesterol exists only as a minor component. Sitosterol and stigmasterol are the most abundant and widely distributed plant sterols, while ergosterol is the major occurring sterol in fungus and yeast. The plant sterols are characterized by an additional alkyl group at C-24 on the cholesterol nucleus with either $\alpha$ or $\beta$ chirality. Sterols with methylene and ethylidene substitutes are also found in plants (e.g., 24-methylene cholesterol, fucosterol). The other major characteristics of plant sterols are the presence of additional double bonds in the side chain, as in porifeasterol, cyclosadol, and closterol.

Despite the diversity of plant sterols and sterols of invertebrates, cholesterol is considered the most important sterol. Cholesterol is an important structural component of cell membranes and is also the precursor of bile acids and steroid hormones [28]. Cholesterol and its metabolism are of importance in human disease. Abnormalities in the biosynthesis or metabolism of cholesterol and bile acid are associated with cardiovascular disease and gallstone formation [29,30]. Our discussion will mainly focus on cholesterol and its metabolites, with a brief comparison of the biosynthesis of cholesterol and plant sterols (see Sec. II.B.2). The biosynthesis of plant sterols and sterols of invertebrates was reviewed by Goodwin [31] and Ikekawa [32].

The chemistry of sterols encompasses a large amount of knowledge relating to the chemical properties, chemical synthesis, and analysis of sterols. A detailed discussion on all these topics is impossible in one chapter. We consider the analysis of sterols to be of primary interest, and therefore our treatment of the chemistry of

Figure 1  Examples of naturally occurring sterols.

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sterols is confined to the isolation, purification, and characterization of sterols from various sources. Readers interested in the chemical reactions and total syntheses of sterols may refer to the monographs in these areas [33–35].

B. Biosynthetic Origins of Sterols

1. Cholesterol Biosynthesis

Cholesterol is the principal mammalian sterol and the steroid that modulates the fluidity of eukaryotic membranes. Cholesterol is also the precursor of steroid hormones such as progesterone, testosterone, estradiol, cortisol, and vitamin D. The elucidation of the cholesterol biosynthesis pathway has challenged the ingenuity of chemists for many years. The early work of Konrad Bloch in the 1940s showed that cholesterol is synthesized from acetyl coenzyme A (acetyl CoA) [36]. Acetate isotopically labeled in its carbon atoms was prepared and fed to rats. The cholesterol that was synthesized by these rats contained the isotopic label, which showed that acetate is a precursor of cholesterol. In fact, all 27 carbon atoms of cholesterol are derived from acetyl CoA. Since then, many chemists have put forward enormous efforts to elucidate this biosynthetic pathway, and this work has yielded our present detailed knowledge of sterol biosynthesis. This outstanding scientific endeavor was recognized by the awarding of several Nobel prizes to investigators in research areas related to sterol [1].

The cholesterol biosynthetic pathway can be generally divided into four stages: (a) the formation of mevalonic acid from three molecules of acetyl CoA; (b) the biosynthesis of squalene from six molecules and mevalonic acid through a series of phosphorylated intermediates; (c) the biosynthesis of lanosterol from squalene via cyclization of 2,3-epoxysqualene; and (d) the modification of lanosterol to produce cholesterol.

The first stage in the synthesis of cholesterol is the formation of mevalonic acid and isopentyl pyrophosphate from acetyl CoA. Three molecules of acetyl CoA are combined to produce mevalonic acid as shown in Scheme 1. The first step of this synthesis is catalyzed by a thiolase enzyme and results in the production of acetoacetyl CoA, which is then combined with third molecule of acetyl CoA by the action of 3-hydroxy-3-methylglutaryl CoA (HMG-CoA) is its cleavage to acetyl CoA and acetoacetate. Acetoacetate is further reduced to β-3-hydroxybutyrate in the mitochondrial matrix. Since it is a β-keto acid, acetoacetate also undergoes a slow, spontaneous decarboxylation to acetone. Acetoacetate, β-3-hydroxybutyrate, and acetone, sometime referred to as ketone bodies, occur in fasting or diabetic individuals. Alternatively, HMG-CoA can be reduced to mevalonate and is present in both the cytosol and the mitochondria of liver cells. The mitochondrial pool of this intermediate is mainly a precursor of ketone bodies, whereas the cytoplasmic pool gives rise to mevalonate for the biosynthesis of cholesterol.

The reduction of HMG-CoA to give the mevalonic acid is catalyzed by a microsomal enzyme, HMG-CoA reductase, which is of prime importance in the control of cholesterol biosynthesis. The biomedical reduction of HMG-CoA is an essential step in cholesterol biosynthesis. The reduction of HMG-CoA is irreversible and proceeds in two steps, each requiring NADPH as the reducing reagent. A hemithioacetal derivative of mevalonic acid is considered to be an intermediate. The concentration of HMG-CoA reductase is determined by rates of its synthesis and
Scheme 1 Synthesis of mevalonic acid from acetyl CoA.

degradation, which are in turn regulated by the amount of cholesterol in the cell. Cholesterol content is influenced by the rate of biosynthesis, dietary uptake, and a lipoprotein system that traffics in the intercellular movement of cholesterol. During growth, cholesterol is mainly incorporated into the cell membrane. However, in homeostasis cholesterol is mainly converted to bile acids and is transported to other tissues via low density lipoprotein (LDL). High density lipoprotein (HDL) also serves as a cholesterol carrier, which carries cholesterol from peripheral tissues to the liver. The major metabolic route of cholesterol is its conversion to bile acids and neutral sterols, which are excreted from the liver via the bile. Kandutsch and Chen and others have shown that oxysterols regulate the biosynthesis of HMG-CoA reductase as well as its digression, which controls cholesterol biosynthesis [37]; the regulation of HMG-CoA reductase by oxysterols is discussed in more detail in a later section. A number of substrate analogs have been tested for their inhibition of HMG-CoA reductase. Some of them (e.g., compactin and melinolin) were found to be very effective in treating hypocholesterol diseases [38,39].

The coupling of six molecules of mevalonic acid to produce squalene proceeds through a series of phosphorylated compounds. Mevalonate is first phosphorylated by mevalonic kinase to form a 5-phosphomevalonate, which serves as the substrate for the second phosphorylation to form 5-pyrophosphomevalonate (Scheme 2). There is then a concerted decarboxylation and loss of a tertiary hydroxy group from 5-pyrophosphomevalonate to form 3-isopentyl pyrophosphate, and in each step one molecule of ATP must be consumed. 3-Isopentyl pyrophosphate is regarded as the basic biological “isoprene unit” from which all isoprenoid compounds are elaborated. Squalene is synthesized from isopentyl pyrophosphate by sequence coupling reactions. This stage in the cholesterol biosynthesis starts with the isomerization of isopentyl pyrophosphate to dimethylallyl pyrophosphate. The coupling reaction shown in Scheme 2 is catalyzed by a soluble sulfhydryl enzyme, isopentyl pyrophosphate–dimethylallyl pyrophosphate isomerase. The coupling of these two isomeric
Scheme 2 Synthesis of isopentenyl pyrophosphate, the biological "isoprene unit," and dimethylallyl pyrophosphate.

C5 units yields geranyl pyrophosphate, which is catalyzed by geranyl pyrophosphate synthetase (Scheme 3). This reaction proceeds by the head-to-tail joining of isopentyl pyrophosphate to dimethylallyl pyrophosphate. A new carbon–carbon bond is formed between the C-1 of dimethylallyl pyrophosphate and C-4 of isopentyl pyrophosphate. Consequently, geranyl pyrophosphate can couple in a similar manner with a second molecule of isopentyl pyrophosphate to produce farnesyl pyrophosphate (C15 structure). The last step in the synthesis of squalene is a reductive condensation of two molecules of farnesyl pyrophosphate (Scheme 4). This step is actually a two-step sequence, catalyzed by squalene synthetase. In the first reaction, presqualene pyrophosphate is produced by a tail-to-tail coupling of two farnesyl pyrophosphate molecules. In the following conversion of presqualene pyrophosphate to squalene, the cyclopropane ring of presqualene pyrophosphate is opened with a loss of the pyrophosphate moiety. A molecule of NADPH is required in the second conversion.

The third stage of cholesterol biosynthesis is the cyclization of squalene to lanosterol (Scheme 5). Squalene cyclization proceeds in two steps requiring, molecular oxygen, NADPH, squalene epoxidase, and 2,3-oxidosqualene–sterol cyclase. The first step is the epoxidation of squalene to form 2,3-oxidosqualene–sterol cyclase. The 2,3-oxidosqualene is oriented as a chair–boat–chair–boat conformation in the enzyme active center. The acid-catalyzed epoxide ring opening initiates the cyclization to produce a tetracyclic protosterol cation. This is followed by a series of concerted 1,2-trans migrations of hydrogen and methyl groups to produce lanosterol.

The last stage of cholesterol biosynthesis is the metabolism of lanosterol to cholesterol. Scheme 6 gives the general biosynthetic pathway from lanosterol to cholesterol.
Scheme 3 Synthesis of farnesyl pyrophosphate from the biological “isoprenyl unit.”

cholesterol. The C-14 methyl group is first oxidized to an aldehyde, and removed as formic acid. The oxidation of the C-4α methyl group leads to an intermediate, 3-oxo-4α-carboxylic acid, which undergoes a decarboxylation to form 3-oxo-4β-methylsterol. This compound is then reduced by an NADPH-dependent microsomal 3-oxosteroid reductase to produce 3β-hydroxy-4α-methyl sterol, which undergoes a similar series of reactions to produce a 4,4-dimethylsterol. In animal tissues, C-14 demethylation and the subsequent double-bond modification are independent of the reduction of the Δ24 double bond. Desmosterol (cholesta-5,24-dien-3β-ol) is found in animal tissues and can serve as a cholesterol precursor. The double-bond isomerization of 8 to 5 involves the pathway $\Delta^8 \rightarrow \Delta^7 \rightarrow \Delta^{5,7} \rightarrow \Delta^5$.

2. Biosynthesis of Plant Sterols

In animals, 2,3-oxidosqualene is first converted to lanosterol through a concerted cyclization reaction. This reaction also occurs in yeast. However, in higher plants and algae the first cyclic product is cycloartenol (Scheme 7). The cyclization intermediate, tetracyclic protosterol cation, undergoes a different series of concerted 1,2-trans migrations of hydrogen and methyl groups. Instead of the 8,9 double bond, a stabilized C-9 cation intermediate is formed. Following a trans elimination of enzyme-X$^- $ and H$^+$ from C-19, with the concomitant formation of the 9,19-cyclopropane ring, cycloartenol is formed. A nearby α-face nucleophile from the enzyme is necessary to stabilize the C-9 cation and allow the final step to be a trans elimination.
Scheme 4  Synthesis of squalene from the coupling of two molecules of farnesyl pyrophosphate.

according to the isoprene rule. The biosynthesis pathway from acetyl CoA to 2,3-oxoisqualene in plants is the same as that in animals (see detailed discussion of the biosynthesis of cholesterol, Sec. II.B.1).

The conversion of cycloartenol to other plant sterols can be generally divided into three steps, which are the alkylation of the side chain at C-24, demethylation

Scheme 5  Cyclization of squalene.
of the C-4 and C-14 methyl groups, and double-bond manipulation. Alkylation in the formation of plant sterols involves methylation at C-24 with S-adenosylmethionine to produce C28 sterols. The further methylation of a C-24 methylene substrate yields C-24 ethyl sterols. The details of the mechanism of demethylation and double-bond manipulation in plants are not clear, but it is highly likely to be very similar to that in animals. In plants, C-4 methyl groups are removed before the methyl group at C-14, whereas in animals it is the other way around. Sterols found in plants are very diversified. The structural features of major plant sterols are depicted in Figure 2.

C. Regulation of Sterol Biosynthesis in Animals

Sterol biosynthesis in mammalian systems has been intensely studied for several decades. Interest in the cholesterol biosynthesis pathway increased following clinical observations that the incidence of cardiovascular disease is greater in individuals with levels of serum cholesterol higher than average. More recently, the results of numerous clinical studies have indicated that lowering serum cholesterol levels may
Scheme 7  Cyclization of squalene to cycloartenol.

Figure 2  Examples of plant sterols.
reduce the risk of coronary heart disease and promote the regression of atherosclerotic lesions [40,41]. The total exchangeable cholesterol of the human body is estimated at 60 g for a 60-kg man. The cholesterol turnover rate is in the order of 0.8–1.4 g/day. Both the dietary uptake and the biosynthesis contribute significantly to body cholesterol. In Western countries the daily ingestion of cholesterol ranges from 0.5 to 3.0 g, although only a portion of this sterol is absorbed from the intestine. The absorption of dietary cholesterol ranges from 25% (high dietary sterol intake) to 50% (low dietary sterol intake). The total body cholesterol level is determined by the interaction of dietary cholesterol, the excretion of cholesterol and bile acid, and the biosynthesis of cholesterol in tissue.

The major site of cholesterol synthesis in mammals is the liver. Appreciable amounts of cholesterol are also formed by the intestine. The rate of cholesterol formation by these organs is highly responsive to the amount of cholesterol absorbed from dietary sources. This feedback regulation is mediated by changes in the activity of HMG-CoA reductase. As discussed in connection with pathway for the biosynthesis of cholesterol, this enzyme catalyzes the formation of mevalonate, which is the committed step in cholesterol biosynthesis. Dietary cholesterol suppresses cholesterol biosynthesis in these organs through the regulation of HMC-CoA reductase activity. In 1974, Kandutsch and Chen observed that highly purified cholesterol (in contrast to crude cholesterol) is rather ineffective in lowering HMG-CoA reductase activity in culture cells [37]. This perception led to the recognition that oxidized derivatives of sterols (oxysterols), rather than cholesterol, may function as the natural regulators of HMG-CoA reductase activity. Furthermore, oxysterols display a high degree of versatility ranging from substrates in sterol biosynthesis to regulators of gene expression to cellular transporters.

Cholesterol, triacylglycerols, and other lipids are transported in body fluids to specific targets by lipoproteins. A lipoprotein is a particle consisting of a core of hydrophobic lipids surrounded by a shell of polar lipids and apoproteins. Lipoproteins are classified according to their densities. LDL, the major carrier of cholesterol in blood, has a diameter of 22 nm and a mass of about 3 × 10^6 daltons (Da). LDL is composed of globular particles, with lipid constituting about 75% of the weight and protein (apoprotein B) the remainder. Cholesterol esters (about 1500 molecules) are located at the core, which is surrounded by a more polar layer of phospholipids and free cholesterol. The shell of LDL contains a single copy of apoprotein B-100, a very large protein (514 kDa). The major functions of LDL are to transport cholesterol to peripheral tissues and to regulate de novo cholesterol synthesis at these sites.

As we discussed above, the major site of cholesterol biosynthesis is the liver. The mode of control in the liver has also been discussed: dietary cholesterol (possibly oxysterols) reduces the activity and amount of HMG-CoA reductase, the enzyme catalyzing the committed step of cholesterol biosynthesis. In some tissues, such as adrenal gland, spleen, lung, and kidney, biosynthesis contributes only a relatively small proportion of the total tissue cholesterol, with the bulk being derived by uptake from LDL in the blood. Investigation upon the interaction of plasma LDL with specific receptors on the surface of some nonhepatic cells has led to a new understanding of the mechanisms of cellular regulation of cholesterol uptake, storage, and biosynthesis in peripheral tissues.

Michael Brown and Joseph Goldstein did pioneering work concerning the control of cholesterol metabolism in nonhepatic cells based on studies of cultured human
fibroblasts [42,43]. In general, cells outside the liver and intestine obtain cholesterol from the plasma rather than by synthesizing them de novo. LDL, the primary source of cholesterol, is first bound to a specific high affinity receptor on the cell surface; endocytosis then transfers it to internal lysosomes, where the LDL cholesteryl ester and protein are hydrolyzed. The released cholesterol suppresses the transcription of the gene from HMG-CoA reductase, hence blocking de novo synthesis of cholesterol. In the meantime, the LDL receptor itself is subject to feedback regulation. The raised cholesterol concentration also suppresses new LDL receptor synthesis. So the uptake of additional cholesterol from plasma LDL is blocked. After the drop of HMG-CoA reductase activity, there is a reciprocal increase in the microsomal acyl CoA-cholesterol acyltransferase (ACAT), with the result that the excess free cholesterol is reesterified for storage. Also, the reduction in the rate of cholesterol biosynthesis, which is attributed to uptake of LDL cholesterol by cells, may in fact be due to the presence of a small amount of oxygenated sterol in the LDL [44]. Hydroxylated sterols are known to be far more potent inhibitors of cholesterol biosynthesis and microsomal HMG-CoA reductase activity than is pure cholesterol [45,46]. The development of the hypothesis that oxysterols are regulators of cholesterol biosynthesis has attracted much attention. A comprehensive review has been published by George Schroepfer, Jr. [47a]. This work could lead to the development of new drugs for the treatment of hypocholesterolemia diseases [47b].

D. Cholesterol Metabolism

In mammals, cholesterol is metabolized into three major classes of metabolic products: (a) the C18, C19, and C21 steroid hormones and vitamin D; (b) the fecal neutral sterols, such as 5α-cholestan-3β-ol and 5β-cholestan-3β-ol; and (c) the C24 bile acids. Only small amounts of cholesterol are metabolized to steroid hormones and vitamin D. These metabolites are very important physiologically. A detailed discussion of steroid hormones is beyond the scope of this chapter. Vitamin D, also considered a steroid hormone, is discussed individually (see Sec. II.E). The neutral sterols and bile acids are quantitatively the most important excretory metabolites of cholesterol.

The fecal excretion of neutral sterols in humans is estimated to range from 0.5 to 0.7 g/day. These sterols are complex mixtures of cholesterol, 5α-cholestan-3β-ol, 5β-cholestan-3β-ol, cholest-4-en-3-one, and a number of cholesterol precursor sterols. The major sterol, 5β-cholestan-3β-ol, is found in the feces as a microbial transformation product of cholesterol.

The principal C24 bile acids are cholic acid and chenodeoxycholic acid. The conversion of cholesterol to bile acids takes place in the liver. These bile acids are conjugated with either glycine or taurine to produce bile salts. The bile salts produced in the liver are secreted into the bile and enter the small intestine, where they facilitate lipid and fat absorption. Most bile acids are reabsorbed from the intestine and pass back to the liver and the enterohepatic circulation. The excretion of bile acids in the feces is estimated to range from 0.4 to 0.6 g/day.

The metabolic pathway of cholesterol to bile acids has been studied for many years. Recent advances in oxysterol syntheses have aided the study of this metabolic pathway [48a–c]. Several reviews are available describing the formation of bile acids from cholesterol [49a,49b]. There are three general stages in the biotransformation
of cholesterol to bile acids (Scheme 8). The first stage is the hydroxylation of cholesterol at the 7α position to form cholest-5-ene-3β,7α-diol. Elucidation of the role of LXR receptors has furthered our knowledge of 7α-hydroxylase and the role of oxysterols in sterol metabolism [48d]. In the second stage, cholest-5-ene-3β,7α-diol is first oxidized to 7α-hydroxycholest-5-en-3-one, which is isomerized to 7α-hydroxycholest-4-en-3-one. Further enzymatic transformation leads to 5β-cholesta-3β,7α-diol and 5β-cholesta-3β,7α,12α-triol. The third stage is the degradation of the hydrocarbon side chain, which is less well understood. However, in cholic acid formation it is generally considered to commence when the steroid ring modifications have been completed. The side chain oxidation begins at the C-26 position; 3β, 7α,12α-trihydroxy-5β-cholestan-26-oic acid is an important intermediate. The removal of the three terminal atoms is believed to proceed by a β-oxidation mechanism analogous to that occurring in fatty acid catabolism.

Scheme 8  Biosynthesis of cholic acid from cholesterol.
E. The Chemistry of Vitamin D and Related Sterols

The discovery of vitamin D dates back to the 1930s, following studies of rickets, a well-known disease resulting from deficiency of vitamin D [50]. Vitamin D has basically two functions in mammals: to stimulate the intestinal absorption of calcium and to metabolize bone calcium. A deficiency of vitamin D results in rickets in young growing animals and osteomalacia in adult animals. In both cases, the collagen fibrils are soft and pliable and are unable to carry out the structural role of the skeleton. As a result, bones become bent and twisted under the stress of the body’s weight and muscle function.

Vitamin D is obtained from dietary uptake or via biosynthesis in the skin by means of the ultraviolet irradiation of 7-dehydrocholesterol. The UV irradiation of 7-dehydrocholesterol first produces provitamin D₃, which results from a rupture in the 9–10 bond followed by a 5,7-sigmatropic shift (Scheme 9). Provitamin then undergoes the thermally dependent isomerization to vitamin D₃ in liver, and further is metabolized to 1,25-dihydroxyvitamin D₃, which is 10 times more active than vitamin D₃, whereas 25-hydroxyvitamin D₃ (Scheme 10) is approximately twice as active as vitamin D₃.

The most important nutritional forms of vitamin D are shown in Figure 3. Of these structures, the two most important are vitamin D₂ and vitamin D₃. These two forms of vitamin D are prepared from their respective 5,7-diene sterols. Vitamins D₄, D₅, and D₆ have also been prepared chemically, but they have much lower biological activity than vitamins D₂ and D₃. Also, many analogs of vitamin D metabolites have been synthesized. Some of these compounds exhibit similar vitamin D hormone responses and have found use in the treatment of vitamin D deficiency diseases (Fig. 4). Recently, vitamin D metabolites were found to be potent inducers of cancer cells, which make this steroid hormone and its analogs (biosynthetic inhibitors) potential candidates for the treatment of cancers and other diseases [51].

Scheme 9  Photochemical synthesis of vitamin D₃.
The most interesting analogs possessing hormonal activity are the 26,26,26,27,27,27-hexafluoro-1,25-(OH)2D3 and 24,24-F2-1,25(OH)2D3, which possess all responses to the vitamin D hormone but are 10–100 times more active than the native hormone. The fluoro groups on the side chain block the metabolism of these compounds.

All vitamin D compounds possess a common triene structure. Thus, it is not surprising that they have the same ultraviolet maximum at 265 nm, a minimum of...
228 nm, and a molar extinction coefficient of 18,200. Since they possess intense ultraviolet absorption, they are labile to light-induced isomerization. In addition, the triene system is easily protonated, resulting in isomerization that produces isotachysterol, which is essentially devoid of biological activity. The lability of the triene structure has markedly limited the chemical approaches to modification of this molecule. There is a great deal of chemistry relating to the chemical properties and syntheses of vitamin D compounds. The detailed chemistry and chemical synthesis of the D vitamins are beyond the scope of this chapter, and interested readers are referred to reviews in this important area [52,53].

F. Analysis of Sterols

1. Extraction of Sterols

To analyze the sterols in specific biological tissues, sterols are first extracted from these tissues with organic solvents. The choice of an extraction technique is often determined by the nature of the source and the amount of information the investigator chooses to obtain concerning the forms of sterols present as free, glycosylated, and/or esterified through the 3-hydroxy group. Different extraction procedures may vary dramatically depending on the extraction efficiency required for different classes of sterols [54,55]. Irrespective of the nature of the source, one of four methods of
sample preparation is usually employed. Samples can be extracted directly, with little or no preparation; after drying and powdering; after homogenization of the fresh materials; or after freeze-drying or fresh-freezing, followed by powdering, sonication, or homogenization. Extraction procedures vary. The analyst may simply mix the prepared material with the extraction solvent (the most frequently used solvents include mixtures of chloroform and methanol, or dichloromethane and acetone) for a short time (0.5–1 hour) and separate the organic solvent phase from the aqueous phases and debris by centrifugation. Another common procedure is extraction from a homogenized material by means of a refluxing solvent in a Soxhlet apparatus for 18 hours or with a boiling solvent for 1 hour. To obtain a total lipid extraction, saponification under basic (i.e., in 10% KOH in 95% ethanol) or acidic conditions is usually conducted prior to the organic solvent extraction. Most of the glycosylated sterols and some esterified sterols cannot be easily extracted into organic solvents without the hydrolysis step.

Many oxysterols contain functional groups (e.g., epoxides and ketones) that may be sensitive to high concentration of acids or bases. Epoxides may undergo nucleophilic attack by strong bases (e.g., NaOH and KOH), followed by ring opening. Moreover, treatment with strong acids can result in ring opening to form to alcohols, alkenes, and ketones. The hydrolysis of cholesterol epoxides under mildly acidic conditions has been studied [56]. Modified procedures are available for the isolation of steroidal epoxides from tissues and cultured cells by saponification and/or extraction [57–59]. Ketones are known to form enolates under the influence of strong bases, which may then form condensation products of higher molecular weight [60]. To circumvent these potential problems, procedures using different extraction techniques are sometimes preferred to a saponification followed by extraction. Mild methods for the removal of the ester function without ketone enolization include extraction by means of sodium or potassium carbonate in heated aqueous solutions of methanol or ethanol. The addition of tetrahydrofuran to these mixtures has been found to significantly increase the solubility of the more polar oxysterols [61].

There are not many studies on the efficiency of various extraction methods. Most extraction procedures were designed to compare the extraction of lipids from cells or tissues of a single source and have been applied subsequently to plants and animals of various types. The errors in the quantitative analysis of sterols, which are probably introduced in the extraction steps, could be eliminated by using in situ labeling of key sterols. Sterols labeled with deuterium and 14C have been used to monitor the extraction recovery in human plasma oxysterol analysis.

2. Isolation of Sterols

Conventional column chromatography, with ordinary phase (silica gel or alumina oxide), reversed phase, and argentation stationary phase, is still the most important method for the isolation and purification of sterols, especially if the total lipid extraction is complex and high in weight (>200 mg) [62]. Chromatographic methods with an organic phase involve the binding of a substrate to the surface of a stationary polar phase through hydrogen bonding and dipole–dipole interaction. A solvent gradient with increasing polarity is used to elute the substrate from the stationary phase. The order of substrate movement will be alkyl > ketone > hindered alcohol > unhindered alcohol. The elution profile is routinely monitored by GC or TLC. Reversed phase column chromatography involves the use of lipophilic dextran (Sephadex LH-
Argentation column chromatography is a very powerful chromatographic method in the separation of different alkene isomers. The argentation stationary phase is typically made by mixing AgNO₃ solution (10 g of AgNO₃ in 10 mL of water) and silica gel or aluminum oxide (90 g of stationary phase in 200 mL of acetone). After acetone has been evaporated at a moderate temperature (<35°C) under vacuum, the resulting argentation stationary phase is dried under vacuum at room temperature until a constant weight is obtained. Usually argentation chromatography is suitable only for the separation of sterol acetates. For free sterol, on-column decomposition of substrate was observed. Compared to the conditions of normal adsorption column chromatography, a slightly polar solvent system is usually needed to elute the substrate from the column. For a preliminary clean-up step, good normal silica gel column chromatography running under the gravity will give good results. Column chromatography with fine silica gel and/or aluminum oxide (230–400 mesh) running under medium pressure (10–100 psi, MPLC) could be used in the separation of individual sterols and sterol subclasses.

TLC is primarily used as an analytical method to detect compounds. Also, small samples (<20 mg) can be separated by means of preparative TLC. For an analytical TLC plate, the thickness of the stationary phase (usually silica gel) is less than 0.25 mm; for preparative TLC a much thicker stationary phase (>0.5 mm) is required. No more than 1 mg of substrate should be loaded on 1-cm analytical plates, and 2 mg on preparative plates. Otherwise overloading will prevent the achievement of a good separation. Nondestructive detection methods (such as iodine and UV detection) are usually used to locate individual compounds. The silica gel containing pure individual compounds is scraped from the plates and eluted with polar solvents.

HPLC, adsorption or reversed phase, is becoming the most commonly used technique for the separation of individual sterols from subclass fractions [62,63]. In adsorption HPLC, microspheres of silica (silicic acid such as µ-Porasil) are employed. As in adsorption, open column, and thin-layer chromatography, this method is characterized by hydrogen bonding and other electronic attractions between the sterol and stationary phase. The smaller diameter (3–5 μm) and high porosity of the microspheres create a very large effective surface area and allow the simultaneous analysis of many more theoretical plates than the other systems can handle. TLC solvent systems can be used directly for adsorption HPLC. In general, the solvents that give best separations are binary (or trinary) systems of mostly low to moderate polarity solvents with a small amount of a strongly polar solvent such as hexane/benzene (9:1) or dichloromethane–n-hexane-ethyl acetate (94:5:1).

In reversed phase HPLC (RP-HPLC) systems, alkyl groups at C-8 or C-18 have been chemically bonded to microspheres of silica. These small diameters (2–5 μm), highly porous particles give the stationary phases very large effective surface areas for the interaction of the sterols with the bonded alkyl groups. This large area, in turn, makes possible rapid equilibration of the sterol between the stationary phase and an appropriate mobile phase, yielding columns with as many as 10⁴ theoretical plates. A variety of polar mobile phases have been employed in RP-HPLC [e.g., 100% acetonitrile, acetonitrile–water mixtures (0–20%), and methanol–water mixtures (2–20% water)]. Samples are usually injected onto the analytical column.
at levels of 1–100 μg per component (up to 1.0 mg total) for routine separations. Retention volumes of eluted sterols are usually expressed relative to cholesterol.

Gas–liquid chromatography (GLC) is most frequently used as an analytical technique to monitor fractions during the isolation and separation of sterols; however, this technique can also be used in preparative separations. The use of GLC as an analytical and preparative technique for sterols and related steroids has been discussed in numerous reviews [64,65]. The separation of sterols in gas–liquid systems depends on the polarity and molecular weight (frequently correlating with size and volume) of the molecule.

3. Characterization of Sterols

GLC is most frequently used to quantitate sterols in extracts, subfraction mixtures, and isolated sterol fractions. Selectivity is provided by gas–liquid partitioning. In general, three different methods of detection have been employed for quantitation: flame ionization detection (FID), electron capture detection (ECD), and mass detection (MD) [66,67]. However, FID is by far the most commonly employed method because it is relatively insensitive to temperature changes during analysis and to minor structural differences in sterols and related steroids, and because it has a large linear mass range of response. For quantitative analysis it is necessary to determine the linear range or response to a sterol standard (e.g., cholesterol or 24-cholestanol) for each FID and accompanying chromatographic system. For sample analysis, dried fractions are routinely weighed and dissolved in a known amount of solvent to give a mass-to-volume ratio within the linear range of the detector, whereupon the samples are injected onto the column.

Quantification of sterols in extracts, subclass fractions, or isolated sterol fractions by HPLC is somewhat limited. The selectivity provided by either adsorption or reversed phase chromatography is in many cases greater than that provided by GLC. Most frequently, UV detectors are employed and are set at end-adsorption wavelengths [68]. These detectors have limited sensitivity, with monochromatic detectors being most sensitive. Additionally, UV detectors cannot be considered universal sterol detectors. Most sterols differ significantly in their UV absorption properties, even in end-absorption regions. Thus, it would not be possible to select a single wavelength for the quantitation of complex sterol mixtures. Therefore, HPLC coupled to variable-wavelength detectors or multiwavelength detectors can be used to quantitate specific sterols in a mixture if the compound in question has a unique absorption spectrum relative to other members of the mixture.

For a full discussion of the identification of sterols with NMR [69,70], mass [71], UV [72], infrared, and X-ray [73] spectrographic techniques, the reader is referred to recent comprehensive reviews in this area.

REFERENCES


72. P. Acuna-Johnson and A. C. Oehlschlager. Identification of sterols and biologically significant steroids by UV and IR spectroscopy. In: Analysis of Sterols and Other

I. INTRODUCTION

Lipids are among the major components of food of plant and animal origin. There is no precise definition available for the term lipid; however, it usually includes a broad category of compounds that have some common properties and compositional similarities. Lipids are materials that are sparingly soluble or insoluble in water, but soluble in selected organic solvents such as benzene, chloroform, diethyl ether, hexane, and methanol. Together with carbohydrates and proteins, lipids constitute the principal structural components of tissues. However, the common and unique features of lipids relate to their solubility rather than their structural characteristics [1]. Many classification systems have been proposed for lipids. From the nutrition point of view, according to the National Academy of Sciences’ report on nutrition labeling, fats and oils are defined as the complex organic molecules that are formed by combining three fatty acid molecules with one molecule of glycerol [2]. As indicated in Table 1 [3–5], lipids are generally classified as simple and compound (complex) or derived lipids according to the Bloor [3] classification.

Foods contain any or all of these lipid compounds; however, triacylglycerols (TAGs) and phospholipids (PLs) are the most abundant and important ones. Liquid TAGs at room temperature are referred to as oils, and are generally of plant or marine origin (e.g., vegetable and marine oils). Solid TAG at room temperature are termed fats, which are generally of animal origin (e.g., lard and tallow).

Accurate and precise analysis of lipids in foods is important for determining constituting components and nutritive value, standardizing identity and uniformity,
Table 1 General Classification of Lipids

<table>
<thead>
<tr>
<th>Simple lipids: Compounds with two types of structural moieties</th>
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</thead>
<tbody>
<tr>
<td>Glyceryl esters: esters of glycerol and fatty acids (e.g., triacylglycerols, partial acylglycerols)</td>
</tr>
<tr>
<td>Cholesteryl esters: esters of cholesterol and fatty acids</td>
</tr>
<tr>
<td>Waxes: true waxes are esters of long chain alcohols and fatty acids; esters of vitamins A and D are also included</td>
</tr>
<tr>
<td>Ceramides: amides of fatty acids with long chain di- or trihydroxy bases containing 12–22 carbon atoms in the aliphatic chain (e.g., sphingosine)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Complex lipids: Compounds with more than two types of structural moieties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phospholipids: glycerol esters of fatty acids, phosphoric acid, and other groups containing nitrogen</td>
</tr>
<tr>
<td>Phosphatidic acid: diacylglycerol esterified to phosphoric acid</td>
</tr>
<tr>
<td>Phosphatidylcholine: phosphatidic acid linked to choline, lecithin</td>
</tr>
<tr>
<td>Phosphatidylethanolamine</td>
</tr>
<tr>
<td>Phosphatidyserine</td>
</tr>
<tr>
<td>Phosphatidylinositol</td>
</tr>
<tr>
<td>Phosphatidyl acylglycerol: more than one glycerol molecule is esterified to phosphoric acid (e.g., cardiolipin, diphosphatidyl acylglycerol)</td>
</tr>
<tr>
<td>Glycoglycerolipids: 1,2-diacylglycerol joined by a glycosidic linkage through position sn-3 to a carbohydrate moiety [e.g., monogalactosyl diacylglycerol, digalactosyl monoacylglycerol, sulfoquinovosyl diacylglycerol (monogalactosyl diacylglycerol at position 6 of the disaccharide moiety is linked by carbon–sulfur bond to a sulfonic acid)]</td>
</tr>
<tr>
<td>Gangliosides: glycolipids that are structurally similar to ceramide polyhexoside and also contain 1–3 sialic acid residues, most contain an amino sugar in addition to the other sugars</td>
</tr>
<tr>
<td>Sphingolipids: derivatives of ceramides</td>
</tr>
<tr>
<td>Sphingomyelins: ceramide phosphorylcholine</td>
</tr>
<tr>
<td>Cerebrosides: ceramide monohexoside (i.e., ceramide linked to a single sugar moiety at the terminal hydroxyl group of the base)</td>
</tr>
<tr>
<td>Ceramide dihexoside: linked to a disaccharide</td>
</tr>
<tr>
<td>Ceramide polyhexoside: linked to a tri- or oligosaccharide</td>
</tr>
<tr>
<td>Cerebroside sulfate: ceramide monohexoside esterified to a sulfate group</td>
</tr>
</tbody>
</table>

| Derived lipids: Compounds that occur as such or released from simple or complex lipids due to hydrolysis (e.g., fatty acids; fatty alcohols; fat-soluble vitamins A, D, E, and K; hydrocarbons; sterols) |

Source: Adapted from Ref. [3–5].
preparing nutritional labeling material, as well as for promoting and understanding the effects of fats and oils on food functionality. At the same time, knowledge about the structural characteristics of lipids may allow development of tailor-made products designed for a particular function or application.

II. EXTRACTION OF LIPIDS FROM FOODS AND BIOLOGICAL MATERIALS

Lipids in nature are associated with other molecules via (a) van der Waals interaction, e.g., interaction of several lipid molecules with proteins; (b) electrostatic and hydrogen bonding, mainly between lipids and proteins; and (c) covalent bonding among lipids, carbohydrates, and proteins. Therefore, to separate and isolate lipids from a complex cellular matrix, different chemical and physical treatments must be administered. Water insolubility is the general property used for the separation of lipids from other cellular components. Complete extraction may require longer extraction time or a series or combination of solvents so that lipids can be solubilized from the matrix.

The existing procedures of lipid extraction from animal or plant tissues usually include several steps: (a) pretreatment of the sample, which includes drying, size reduction, or hydrolysis; (b) homogenization of the tissue in the presence of a solvent; (c) separation of liquid (organic and aqueous) and solid phases; (d) removal of nonlipid contaminants; and (e) removal of solvent and drying of the extract. Standard methods for lipid extraction have been established by the Association of Official Analytical Chemists (AOAC) International for different types of materials/tissues. However, when it comes to practical situations, each case might require modification of the method.

A. Sample Preparation

As with any form of chemical analysis, proper sampling and storage of the samples are essential for obtaining valid results. According to Pomeranz and Meloan [5], an ideal sample should be identical in all of its intrinsic properties to the bulk of the material from which it is taken. In practice, a sample is satisfactory if its properties under investigation correspond to those of the bulk material within the limits set by the nature of the test. Sample preparation for lipid analysis depends on the type of food and the nature of its lipids. Effective analysis calls for a knowledge of the structure, chemistry, and occurrence of principal lipid classes and their constituents. Therefore, it is not possible to devise a single standard method for extraction of all kinds of lipids in different foods.

Extraction of lipids should be performed as soon as possible after the removal of tissues from the living organism so as to minimize any subsequent changes. Immediate extraction is not always possible; however, the samples usually are stored at very low temperatures in sealed containers, under an inert (nitrogen) atmosphere or on dry ice. Yet the freezing process itself may permanently damage the tissues as a result of osmotic shock, which alters the original environment of the tissue lipids and brings them into contact with enzymes from which they are normally protected. Thawing the sample taken from frozen storage before extraction may enhance this deterioration. Therefore, tissue samples should be homogenized and extracted with
solvents without being allowed to thaw [4]. Lipolytic enzymes of animal and plant tissues are usually deactivated irreversibly by homogenization with polar solvents. Use of high temperatures should be avoided; it is also advisable, when possible, to maintain an inert atmosphere during sample preparation and extraction which may minimize oxidation reactions of unsaturated lipids.

B. Pretreatments

1. Drying

Sometimes nonpolar solvents, such as diethyl ether and hexane, do not easily penetrate the moist tissues (>8% moisture); therefore, effective lipid extraction does not occur. Diethyl ether is hygroscopic and becomes saturated with water and thus inefficient for lipid extraction. Therefore, reducing moisture content of the samples may facilitate lipid extraction. Vacuum oven drying at low temperatures or lyophilization is usually recommended. Predrying facilitates the grinding of the sample, enhances extraction, and may break fat-water emulsions to make fat dissolve easily in the organic solvent and helps to free tissue lipids. Drying the samples at elevated temperatures is undesirable because lipids become bound to proteins and carbohydrates, and such bound lipids are not easily extracted with organic solvents [5].

2. Particle Size Reduction

The extraction efficiency of lipids from a dried sample also depends on the size of the particles. Therefore, particle size reduction increases surface area, allowing more intimate contact of the solvent, and enhances lipid extraction (e.g., grinding of oilseeds before lipid extraction). In some cases, homogenizing the sample together with the extracting solvent (or solvent system) is carried out instead of performing these operations separately.

3. Acid/Alkali Hydrolysis

To make lipids more available for the extracting solvent, food matrices are often treated with acid or alkali prior to extraction. Acid or alkali hydrolysis is required to release covalently and ionically bound lipids to proteins and carbohydrates as well as to break emulsified fats. Digestion of the sample with acid (usually 3–6 M HCl) under reflux conditions converts such bound lipids to an easily extractable form. Many dairy products, including butter, cheese, milk, and milk-based products, require alkali pretreatment with ammonia to break emulsified fat, neutralize any acid, and solubilize proteins prior to solvent extraction [6]. Enzymes are also employed to hydrolyze food carbohydrates and proteins (e.g., use of Clarase, a mixture of α-amylase and protease) [2].

C. Lipid Extraction with Solvents

The insolubility of lipids in water makes possible their separation from proteins, carbohydrates, and water in the tissues. Lipids have a wide range of relative hydrophobicity depending on their molecular constituents. In routine food analysis, "fat" content (sometimes called the ether extract, neutral fat, or crude fat) refers to "free" lipid constituents that can be extracted into less polar solvents, such as light petroleum ether or diethyl ether. The "bound" lipid constituents require more
polar solvents, such as alkanols, for their extraction. Therefore, use of a single universal solvent for extraction of lipids from tissues is not possible. During solvent extraction van der Waals and electrostatic interactions as well as hydrogen bonds are broken to different extents; however, covalent bonds remain intact.

Neutral lipids are hydrophobically bound and can be extracted from tissues by nonpolar solvents, whereas polar lipids, which are bound predominantly by electrostatic forces and hydrogen bonding, require polar solvents capable of breaking such bonds. However, less polar neutral lipids, such as TAGs and cholesterol esters, may also be extracted incompletely with nonpolar solvents, probably due to inaccessibility of a significant part of these lipids to the solvents. Lipids that are covalently bound to polypeptide and polysaccharide groups will not be extracted at all by organic solvents and will remain in the nonlipid residue. Therefore, a hydrolysis step may be required to release covalently bound lipids to render them fully extractable.

1. Properties of Solvents and Their Mode of Extraction

The type of solvent and the actual method of lipid extraction depend on both the chemical nature of the sample and the type of lipid extract (e.g., total lipids, surface lipids of leaves) desired. The most important characteristic of the ideal solvent for lipid extraction is the high solubility of lipids coupled with low or no solubility of proteins, amino acids, and carbohydrates. The extracting solvent may also prevent enzymatic hydrolysis of lipids, thus ensuring the absence of side reactions. The solvent should readily penetrate sample particles and should have a relatively low boiling point to evaporate readily without leaving any residues when recovering lipids. The solvents mostly used for isolation of lipids are alcohols (methanol, ethanol, isopropanol, n-butanol), acetone, acetonitrile, ethers (diethyl ether, isopropyl ether, dioxane, tetrahydrofuran), halocarbons (chloroform, dichloromethane), hydrocarbons (hexane, benzene, cyclohexane, isooctane), or their mixtures. Although solvents such as benzene are useful in lipid extraction, it is advisable to look for alternative solvents due to the potential carcinogenicity of such products. Flammability and toxicity of the solvent are also important considerations to minimize potential hazards as well as cost and nonhygroscopicity.

Solubility of lipids in organic solvents is dictated by the proportion of the nonpolar hydrocarbon chain of the fatty acids or other aliphatic moieties and polar functional groups, such as phosphate or sugar moieties, in their molecules. Lipids containing no distinguishable polar groups (e.g., TAGs or cholesterol esters) are highly soluble in hydrocarbon solvents such as hexane, benzene, or cyclohexane and in more polar solvents such as chloroform or diethyl ether, but remain insoluble in polar solvents such as methanol. The solubility of such lipids in alcoholic solvents increases with the chain length of the hydrocarbon moiety of the alcohol; therefore, they are more soluble in ethanol and completely soluble in n-butanol. Similarly, the shorter chain fatty acid residues in the lipids have greater solubility in more polar solvents (e.g., tributyryl is completely soluble in methanol whereas tripalmitin is insoluble). Polar lipids are only sparingly soluble in hydrocarbon solvents unless solubilized by association with other lipids; however, they dissolve readily in more polar solvents, such as methanol, ethanol, or chloroform [4].

2. Extraction Methods with Single Organic Solvent

Diethyl ether and petroleum ether are the most commonly used solvents for extraction of lipids. In addition, hexane and sometimes pentane are preferred to obtain
lipids from oilseeds. Diethyl ether (bp 34.6°C) has a better solvation ability for lipids compared to petroleum ether. Petroleum ether is the low boiling point fraction (bp 35–38°C) of petroleum and mainly contains hexanes and pentanes. It is more hydrophobic than diethyl ether and therefore selective for more hydrophobic lipids [5,7]. The main component (>95%) of dietary lipids are TAGs, while the remaining lipids are mono- and diacylglycerols, phospho- and glycolipids, and sterols. Therefore, nonpolar solvent extractions have been widely employed to extract and determine lipid content of foods. However, oil-soluble flavor, vitamins, and color compounds may also be extracted and determined as lipids when less polar solvents are used.

In determining total lipid content, several equipment and methods have been developed that utilize single-solvent extraction. Among them the gravimetric methods are most commonly used for routine analysis purposes. In gravimetric methods, lipids of the sample are extracted with a suitable solvent continuously, semicontinuously, or discontinuously. The fat content is quantified as weight loss of the sample or by weight of the fat removed. The continuous solvent extraction (e.g., Goldfisch and Foss–Let) gives a continuous flow of boiling solvent to flow over the sample (held in a ceramic thimble) for a long period. This gives a faster and more efficient extraction than semicontinuous methods but may result in incomplete extraction due to channeling. In the semicontinuous solvent extraction (e.g., Soxhlet, Soxtec), the solvent accumulates in the extraction chamber (sample is held in a filter paper thimble) for 5–10 minutes and then siphons back to the boiling flasks. This method requires a longer time than the continuous method, provides a soaking effect for the sample, and does not result in channeling. In the direct or discontinuous solvent extraction, there is no continuous flow of solvent and the sample is extracted with a fixed volume of solvent. After a certain period of time the solvent layer is recovered, and the dissolved fat is isolated by evaporating the organic solvent. Rose-Gottlieb, modified Mojonnier, and Schmid–Boudzynski–Ratzlaff (SBR) methods are examples, and these always include acid or base dissolution of proteins to release lipids [6]. Such procedures sometimes employ a combination extraction with diethyl and petroleum ethers to obtain lipids from dairy products. Use of these solvents may allow extraction of mono-, di-, and triacylglycerols, most of the sterols and glycolipids, but may not remove phospholipids and free fatty acids.

3. Methods Using Organic Solvent Combination

A single nonpolar solvent may not extract the polar lipids from tissues under most circumstances. To ensure a complete and quantitative recovery of tissue lipids, a solvent system composed of varying proportions of polar and nonpolar components may be used. Such a mixture extracts total lipids more exhaustively and the extract is suitable for further lipid characterization. The methods of Folch et al. [8] and Bligh and Dyer [9] are most widely used for total lipid extraction. Use of a polar solvent alone may leave nonpolar lipids in the residue; when lipid-free apoproteins are to be isolated, tissues are defatted with polar solvents only [10]. It is also accepted that the water in tissues or water used to wash lipid extracts markedly alters the properties of organic solvents used for lipid extraction.

Commonly the chloroform–methanol (2:1, v/v) solvent system [8] provides an efficient medium for complete extraction of lipids from animal, plant, or bacterial tissues. The initial solvent system is binary; during the extraction process, it becomes
a ternary system consisting of chloroform, methanol, and water in various proportions, depending on the moisture content of the sample [9]. The method of Bligh and Dyer [9] specifically recognizes the importance of water in the extraction of lipids from most tissues and also plays an important role in purifying the resulting lipid extract. A typical Folch procedure uses a solvent-to-sample ratio of 2:1 (v/w) with a mixture of chloroform and methanol (2:1, v/v) in a two-step extraction. The sample is homogenized with the solvent and the resultant mixture filtered to recover the lipid mixture from the residue. Repeated extractions are usually carried out, separated by washings with fresh solvent mixtures of a similar composition. It is usually accepted that about 95% of tissue lipids are extracted during the first step. In this method, if the initial sample contains a significant amount of water, it may be necessary to perform a preliminary extraction with 1:2 (v/v) chloroform–methanol in order to obtain a one-phase solution. This extract is then diluted with water or a salt solution (0.08% KCl) until the phases separate and the lower phase containing lipids is collected. Bligh and Dyer [9] uses 1:1 (v/v) chloroform–methanol for the first step extraction and the ratio is adjusted to 2:1 (v/v) in the alternate step of extraction and washing. The original procedure of Folch or of Bligh and Dyer uses large amounts of sample (40–100 g) and solvents; therefore, the amounts may be scaled down when a small amount of sample is present or for routine analysis in the laboratory. Hence, Lee and coworkers [11] have described a method that uses the same solvent combination, but in different proportions, based on the anticipated lipid content of the sample. According to this method, chloroform–methanol ratios of 2:1 (v/v) for fatty tissues (>10% lipid) and 1:2 (v/v) for lean (<2%) tissues are recommended.

Folch extraction recovers neutral lipids, diacylglycerophospholipids, and most of the sphingolipids. Lysophospholipids are only partly recovered, and more polar acidic phospholipids and glycolipids may be lost during washing with water. However, both Folch and Bligh–Dyer procedures may fail to transfer all of the lipids to the organic phase. Lysophospholipids, phosphoinositides, and other highly polar lipid substances are selectively lost. According to Christie [5], tissues rich in phosphoinositides should be stored in such a manner as to minimize their enzymatic degradation, and solvent extraction should be performed initially in the presence of CaCl₂. When lysophosphatides are the major component of the tissue extract, it is recommended that acids or inorganic salts be added during extraction with chloroform–methanol, or n-butanol saturated with water. Therefore, specific applications and modifications of the method are required to ensure complete recovery of tissue lipids.

Due to the potential health hazards of chloroform, solvent mixtures containing alkane–alcohol–water mixtures such as hexane and isopropanol, with or without water, have been successfully used to extract tissue [12,13] and fish meal lipids [14]. Hexane–isopropanol (3:2, v/v) [12,14,15], heptane–ethanol–water–sodium dodecylsulfate (1:1:1, v/v/v) [16], methylene chloride–methanol (2:1, v/v) [17,18], and hexane–acetone (1:1, v/v) [19] are such solvent combinations employed to extract lipids from biological materials. Azeotropes of isopropanol have also been used to extract lipids from oilseeds as substitutes for hexane [20–22]. Water-saturated n-butanol [23] has been most effective in extracting lipids from cereals that are rich in starch. This solvent mixture is used extensively for extracting lipids from starchy foods; however, acid hydrolysis might be needed to release bound lipids or inclusion complexes prior to their extraction.
Accelerated solvent extraction (ASE) techniques have recently been introduced. These use classical solvent systems to extract lipids, but under varying extraction parameters such as temperature, pressure, and volume. ASE is also automated [24,25]. The ASE process consumes a much lower solvent volume and time as lipid is extracted at temperatures well above the boiling point of the solvent due to the elevated pressure used in the process. This enhances solubilization and diffusion of lipids from samples into the solvent, significantly shortening the extraction time and solvent consumption. The fat could be extracted with no outflow of solvent (static mode) or allowing fresh solvent to flow continuously through the sample (dynamic mode) during extraction. Under elevated temperature and pressure, dissolved lipids diffuse from the core to the surface of the sample particles and then are transferred to the extraction solvent. Compressed gas then purges the solubilized fat into a collection vessel and can then be quantified gravimetrically [7]. According to Schafer [25], the content of fatty acids of the lipids extracted from muscle matrices using ASE (Dionex 200 or 300 System, chloroform–methanol solvent system) was similar or better in comparison with the conventional Folch extraction. The automated solvent extractors contain microwave moisture analyzer to dry the sample before extraction, redry to remove solvent and moisture, and to determine the percentage of fat as weight loss due to the extraction process [2].

4. Methods Using Nonorganic Solvents

Due to environmental concerns and potential health hazards of organic solvents, nonorganic solvents have become popular. The use of microwave digestion for isolating lipids has recently been reported [26]. It is suggested that microwave energy, by increasing the rotational force on bonds connecting dipolar moieties to adjacent molecules, reduces the energy required to disrupt hydrophobic associations. Hydrogen bonding, and electrostatic forces, thus helping to dissolve all kinds of lipids [26]. Microwave technology has allowed the development of rapid, safe, and cost-effective methods for extracting lipids and does not require that samples be devoid of water [27]. Performance of microwave lipid extraction was qualitatively (all lipid classes) and quantitatively comparable to that of the conventional Folch method for various biological samples [26].

Supercritical fluid extraction (SFE). When carbon dioxide is compressed at a temperature (31.1°C) and pressure (72.9 atm) above its critical point, it doesn’t liquify but attains a dense gaseous state that behaves like a solvent. Thus, it is called supercritical CO₂ (SC-CO₂). Use of SC-CO₂ for lipid extraction significantly reduces the use of organic solvents, avoids waste disposal problems, eliminates the use of potentially toxic and flammable solvents, and reduces the extraction time. Lipids so extracted are not subjected to high temperatures during the extraction process.

 Extraction using SC-CO₂ yields a good recovery of nonpolar lipids including esterified fatty acids, acylglycerols, and unsaponifiable matter. Complex polar lipids are only sparingly soluble in SC-CO₂. The polarity of SC-CO₂ can be varied by using an entrainer such as methanol, ethanol, or even water to improve the extraction of polar lipids [28–31]. This technique has been used for the extraction of lipids from various matrices, including dehydrated foods [32,33], meats [34–36], and fried foods [37]. Particle size also affects lipid recovery because it influences the surface area exposed to SC-CO₂. High moisture content decreases contact between sample and SC-CO₂ as well as the diffusion lipids outside the sample [38]. The extracted
lipids from meat or hydrolytic products from the acid hydrolysis step are allowed to absorb onto a solid phase extraction (SPE) matrix and SC-CO₂ can be used to extract the adsorbed lipids [35]. An increased lipid recovery with decreased moisture content has been demonstrated in wet samples, such as meat [34,39–42]. Therefore, lyophilization is suggested to improve the extraction efficiency of lipids from samples with a high moisture content. The SC-CO₂ extraction is able to recover 97–100% of lipids when compared to the conventional solvent extraction methods [43,44]; no significant differences between fatty acids extracted were observed. Several researchers have shown that supercritical fluid extraction (SFE) could replace solvent extraction methods in a large variety of samples. In fact SFE has recently been included in the recommended methods of the AOAC to extract lipids from oilseeds [45]. The main drawback of SC-CO₂ is equipment cost and the extraction of nonfat materials, such as water [46].

D. Lipid Extraction Without Solvents

Lipid extraction methods are mostly wet extraction procedures that do not use solvents and lipid content is quantified by volumetric means. Such procedures are well utilized in determining fat content of dairy foods, especially fresh milk, and require the use of specifically designed glasswares and equipment.

1. Acid Digestion Methods

Babcock and Gerber methods are classical examples for acid digestion methods. The basic principle of these methods is destabilization and release of fat from the emulsion with a strong acid (e.g., sulfuric). The less dense fat rises in the calibrated neck of the Babcock bottle and the centrifugation step helps the separation. Added sulfuric acid digests proteins, generates heat, and releases fat. The content of fat is measured volumetrically and expressed as weight percent. The modified Babcock method uses an acetic–perchloric acid mixture rather than sulfuric acid and is employed to determine essential oil in flavor extracts and products containing sugar and chocolate. The Gerber method uses a principle similar to that of the Babcock method but utilizes sulfuric acid and pentanol. Pentanol prevents charring of sugar, which can occur with the Babcock method; therefore, the Gerber method could be applied to a wide variety of dairy-based foods [2,6].

2. Detergent Method

The detergent method uses a detergent to form a protein–detergent complex to break up emulsion and release fat. For milk, the anionic detergent dioctyl sodium phosphate is added to disperse the protein layer that stabilizes and liberates fat. Then a strong hydrophilic nonionic polyoxyethylene detergent, sorbitan monolaurate, is added to separate the fat from other food components [5].

3. Physical Methods

External compression forces may be used to release tissue contents and extract lipids, especially from the dry matter. Oilseeds (moisture <5%, oil >30%) are generally subjected to expeller pressing to obtain lipids without using solvents.
E. Removal of Nonlipid Contaminants from Lipid Extracts and Other Practical Considerations

Removal of nonlipid contaminants from the lipid extract is necessary since most of the solvents employed also dissolve significant amounts of oil-soluble flavors, pigments, sugars, amino acids, short chain peptides, inorganic salts, and urea. The nonlipid matter must be removed prior to gravimetric determination of total lipids in order to prevent contamination during subsequent fractionation of the total extract. In chloroform–methanol extract the commonly used method for removing nonlipid contaminants includes washing with water or a diluted KCl solution (0.88%, w/v). Use of salt solution has the advantage of preventing or minimizing the formation of an intermediate phase. When chloroform–methanol (2:1, v/v) is used for extraction of the sample, addition of water or diluted salt solution results in the formation of a two-phase system, i.e., a lower phase consisting of chloroform–methanol–water (86:14:1, v/v/v) and an upper phase consisting of the same, but in the ratio of 3:48:47 (v/v/v). The lower phase composes about two-thirds of the total volume and contains the lipid components, but the upper phase retains the nonlipid contaminants. However, more polar lipids, such as some phospholipids and glycolipids and all gangliosides, may remain in the upper phase [4,47]. Nonlipid contaminants may also be removed partly or completely by evaporation of the lipid extract to dryness in vacuo or under nitrogen and then reextracted with a nonpolar solvent, such as hexane.

In the Bligh and Dyer [9] method the sample is homogenized with chloroform and methanol in such proportions that a miscible system is formed with water in the sample. Dilution of the homogenate with chloroform and water separates it into two layers; the chloroform layer contains all the lipids and the methanol–water layer contains all the nonlipid matter. A purified lipid extract could be obtained by isolating the chloroform layer. Traces of moisture can be removed by passing the chloroform extract through a bed of anhydrous sodium sulfate.

Removal of nonlipid contaminants by liquid–liquid partition chromatography on a dextran gel was introduced by Wells and Dittmer [48]. This is done by passing the crude lipid extract through a Sephadex G-25 column (packed in the upper phase of chloroform–methanol–water, 8:4:3, v/v/v or “Folch wash”). Lipids free of contaminants would be eluted rapidly from the column by employing the lower phase of the Folch wash. Gangliosides and non-lipids are retained in the column and can be recovered by washing with the upper phase of Folch wash and at the same time regenerating the column [49].

Use of predistilled solvents for lipid extraction is advisable since all solvents contain small amounts of lipid contaminants. Use of plastic containers and non-Teflon apparatus should also be avoided as plasticizers may leach out to the lipid extract. To prevent autoxidation of unsaturated lipids it is advisable to add an antioxidant (e.g., BHT) to the solvent (at a level of 50–100 mg/L). Furthermore, extraction should be performed under an inert nitrogen atmosphere, and both tissue and tissue extracts should be stored at −20°C under nitrogen, if possible. Most of the methods described above are suitable for quantifying total lipid content of the sample of interest. When high temperatures are involved in extraction the resulting lipid extract is not suitable for further composition analysis. Folch method–based extraction is usually the preferred procedure to obtain total lipids for further analysis. Lipids are recovered from the chloroform layer by removal of solvent at low tem-
perature under vacuum. Acid hydrolysis also results in decomposition of phospho-
lipids and possibly TAGs to a certain extent [6].

Quantification of lipids from the extracts is mostly carried out as the weight
difference of an aliquot after solvent removal. Removal of the solvent from lipid
extracts should be conducted under vacuum in a rotary evaporator at or near room
temperature. When a large amount of solvent must be evaporated, the solution must
be concentrated and transferred to a small vessel so that the lipids do not dry out as
a thin film over a large area of glass. Lipids should be stored immediately in an inert
nonalcoholic solvent such as chloroform, rather than being allowed to remain in dry
state for long [4].

III. INDIRECT METHODS OF TOTAL LIPID DETERMINATION

Several techniques and instruments have been developed and applied to the indirect
and rapid determination of total lipid content of samples. These methods are really
not lipid extraction methods, but they are gaining popularity because they are rapid
and largely nondestructive. Most of these methods rely on a standard reference pro-
cedure and must be calibrated against a methodology to be validated.

A. Density Measurement

It has been reported that the density of flaxseed is highly correlated ($r = 0.96$) with
its oil content [50]. Thus, measurement of seed density may be used as a means of
screening flax genetic lines for high oil content.

B. Dielectric Method

The dielectric constant of a selected solvent changes when fat is dissolved in it. After
an oilseed sample has been ground with a solvent and the dielectric constant of the
mixture measured, the lipid content is determined from standard charts that show
variation of the dielectric constants of different amounts of lipid in the same solvent
[5]. According to Hunt et al. [51], the amount of induced current and oil content
determined by solvent extraction of soybean are linearly related ($r = 0.98$).

C. Near-Infrared Spectroscopy

The near-infrared (NIR) reflectance is in the range of 14,300–4000 cm$^{-1}$ (700–2500
nm) due to overtone and combination bands of C-H, O-H, and N-H. NIR spectrom-
etry can be used for determination of contents of oil, protein, and moisture and serves
as a very useful tool in the routine analysis of oilseeds.

Rodriguez-Otero et al. [52] have used NIR spectroscopy for measurement of
fat, protein, and total solids of cheese. Lee et al. [53] have used short-wavelength
(700–1100 nm) NIR with a bifurcated fiberoptic probe to estimate the crude lipid
content in the muscle of whole rainbow trout. A very good correlation was observed
between fat content determined by chemical analysis and NIR reflectance spectro-
scopic values obtained for farmed Atlantic salmon fillets [54]. The use of mid-IR
spectroscopy to determine lipid content of milk and dairy products has been de-
scribed by Biggs [55]. Lipids absorb IR energy at the wavelength of 5730 nm, and
the energy absorbed depends on the lipid content of the sample. Quantification is
carried out by the standard curve of the IR absorption and lipid content determined by standard analytical methods [56]. Details about the use of IR spectroscopy for lipid analysis is provided in a later section of this chapter.

D. Low-Resolution Nuclear Magnetic Resonance Spectroscopy

Time domain low-resolution nuclear magnetic resonance (NMR) (referred to as wide-line NMR) and frequency domain NMR could be used to determine the total lipid content of foods. In time domain NMR, signals from the hydrogen nuclei (H or protons) of different food components are distinguished by their different rates of decay or nuclear relaxation. Protons of solid phases relax (signal disappear) quickly, while protons in the liquid phase relax very slowly. Protons of water in the sample relax faster than protons of the lipid. The intensity of the signal is proportional to the number of protons and, therefore, to the hydrogen content. Thus, the intensity of the NMR signal can be converted to oil content of the sample using calibration curves or tables [57–60]. This method can be used to determine the contents of water, oil, and solid–fat and solid-to-liquid ratio of the sample. Time domain NMR has been used to analyze the fat content of foods, including butter, margarine, shortening, chocolate, oilseed, meat, milk and milk powder, and cheese [61–63].

Frequency domain NMR distinguishes food components by resonance frequency (chemical shift) of the peaks in the spectrum. The pattern of oil resonances reflects the degree of unsaturation and other chemical properties [56,61].

E. Turbidimetric/Colorimetric Methods

Haugaard and Pettinati [64] have described a turbidimetric method for rapid determination of lipid in milk. The milk fat is homogenized to obtain uniform globules and the milk proteins are retained with chelating agents such as EDTA. Light transmission of the sample is measured and then converted to the lipid content with the use of a conversion chart.

The lipid content of milk can also be determined by using a colorimetric method [65]. The lipids of milk are allowed to react with an alkaline solution of hydroxamic acid for a specified period. Upon acidification and addition of ferric chloride, a relatively stable chromophore with a maximal absorbance at 540 nm is formed [66].

F. Ultrasonic Method

Fitzgerald et al. [67] have described an ultrasonic method to determine the amount of fat and nonfat solids of liquid milk. The velocity of sound increases or decreases directly with the lipid content above or below a certain critical temperature. This method of fat determination is based on the speed of sound passing through the milk at various temperatures.

G. X-Ray Absorption

It is known that lean meat absorbs more X-ray than high-fat meat [68]. This fact has been used to determine lipid content in meat and meat products using a standard curve of the relationship between X-ray absorption and the lipid content determined by usual solvent extraction methods [5].
IV. ANALYSIS OF LIPID EXTRACTS

Lipid analysis is usually required to determine the composition and structure of the lipid extracted from the sample. Foods must be analyzed to reveal the content and type of saturated and unsaturated lipids as well as their cholesterol content. Such characterization provides information about the caloric value, as well as other properties, including nutritional quality and safety of lipids with respect to their cholesterol and saturated fatty acid contents. In addition, quantification of quality characteristics such as degree of unsaturation, saponification value, refractive index, free fatty acid content, solid fat index, and oxidative stability are required to determine the market value and potential application of fats and oils. Analysis of individual components of lipids is beyond the scope of this chapter and hence is not discussed here.

A. Bulk Oil Properties

The analysis of bulk properties of lipids is primarily important for defining quality characteristics of oils and fats. Therefore, methods applicable to bulk vegetable oils, confectionary fats (e.g., cocoa butter), and table fats (e.g., butter, margarine) are discussed below.

1. Degree of Unsaturation

Iodine value (IV) measures the degree of unsaturation of a lipid and is defined as the number of grams of iodine absorbed by 100 g of lipid. The source of iodine (or other halogen, usually Br₂ and Cl₂) for the reaction is Wijs or Hanus reagent; the reaction involved is essentially a volumetric titration.

In a microanalytical method for determination of the IV of lipids reported by Iskander [69], ethylenic double bonds of the lipid are saturated with bromine vapors, after which the amount of absorbed bromine is determined by neutron activation analysis (NAA).

Determination of IV gives a reasonable quantitation of lipid unsaturation if the double bonds are not conjugated with each other or with a carbonyl oxygen. Furthermore, the determination should be carried out in the absence of light for a given period and with an excess of halogen reagent used [70].

Use of hydrogenation methods to determine degree of unsaturation overcomes the limitations of halogenation methods. Hydrogenation is used to measure the degree of unsaturation of acetylenic or conjugated double bonds. Such fats do not absorb halogen readily; however, the addition of hydrogen to them is considered to be quantitative. This method is essentially a catalytic reaction of heated lipid; the amount of hydrogen absorbed is determined under standard conditions. The results are expressed on a mole basis or on the basis of IV [5].

At the low frequency end of the fingerprint region of IR (1500–900 cm⁻¹) a band due to the CH—CH bending absorption of isolated trans double bonds would be observed. Beyond the isolated trans bond is another group of CH absorption, in this case bending vibrations, including a very strong cis absorption. The combination of cis and trans absorption provides a measure of the total unsaturation or iodine value [71–73].

2. Free Fatty Acid Content

The presence of free fatty acids (FFAs) in an oil is an indication of insufficient processing, lipase activity, or other hydrolytic actions. Classically, the acid value,
which is defined as the number of milligrams of KOH required to neutralize the free acids in 1 g of sample, is a measure of FFA content. FFAs of oils can be determined colorimetrically by dissolving oil in chloroform (or benzene), then allowing the FFAs to react with a cupric acetate solution. The organic solvent turns to a blue color due to the FFA–cupric ion complex, which has a maximum absorbance between 640 and 690 nm [74].

As there is a band attributed to the carboxyl group (COOH) in the center region of the mid-IR spectrum, Fourier transform IR (FTIR) spectroscopy can be used to determine the content of free fatty acids [75,76].

3. Oxidative Stability and Oxidation Products

Owing to their degree of unsaturation, lipids are very susceptible to autoxidation. Autoxidation occurs via a self-sustaining free radical mechanism that produces hydroperoxides (primary products), which in turn undergo scission to form various aldehydes, ketones, alcohols, and hydrocarbons (secondary products). The presence of secondary lipid oxidation products influences the overall quality of a lipid. Methods of determination of oxidative stability and oxidation products are discussed in detail in another chapter.

4. Refractive Index

The refractive index (RI) of an oil is defined as the ratio of the speed of light in vacuum (practically in air) to the speed of light in oil at a specified temperature. This ratio also provides a measure of purity of oils and may be used as a means of identifying them. The RI is measured with a refractometer, usually at 20–25°C for oils and 40°C for solid fats, which generally liquify at 40°C. The RI declines linearly with decreasing IV; thus, it is also used as an index for reporting the degree of hydrogenation of the oil [5].

5. Saponification Value

The saponification value provides an indication of the average molecular weight of lipids. It is defined as the amount of KOH, in milligrams, required to saponify 1 g of fat, i.e., to neutralize the existing FFAs and those liberated from TAG [5].

6. Solid Fat Index

The solid fat index (SFI), an empirical expression of the ratio of liquids in fat at a given temperature, is measured as the change in specific volume with respect to temperature. As a solid fat melts, the volume of the sample increases, and this change is measured by dilatometry. Detection of analysis of phase transformation of fat may also be performed, because lipids expand upon melting and contract upon polymorphic change to a more stable fat [5]. Use of low-resolution pulse NMR and FTIR [77–81] for determination of solid fat content has been detailed in the literature.

B. Chromatographic Procedures for Lipid Characterization

Lipid extracts are complex mixtures of individual classes of compounds and require further separation to pure components if needed. Analysis of chemical components of lipid (e.g., lipid classes, fatty acids, trans fatty acids, sterols, tocopherols, pig-
ments, etc.) primarily involves chromatographic and spectroscopic methods. Usually a combination of separation techniques is used to achieve a high degree of purity of respective lipid components and this could be analytical (for quantitation) or preparative.

The first step in the analyses involves separation of lipids into their various polarity components. It may simply separate the lipid into its polar and nonpolar fractions or may entail analysis of TAG, FFAs, sterols, sterol esters, glycolipids and phospholipids. Traditionally, liquid–liquid extraction, thin-layer chromatography (TLC), or liquid–solid column chromatography have been used for fractionation, cleaning, and concentration of lipid extracts. The most commonly used chromatographic techniques for lipid analysis include column chromatography, gas chromatography (GC), high-performance liquid chromatography (HPLC), supercritical fluid chromatography (SFC), and TLC. Applications of these techniques for the analysis of food lipids are discussed below.

1. Column Chromatography

Lipid extracts are usually fractionated by column chromatography on a preparative scale before subjecting them to detailed analysis. Solid–liquid (adsorption), liquid–liquid (partition), and ion exchange chromatography are among the widely used methods of lipid fractionation. In solid–liquid chromatography, separation is based on partitioning and/or adsorption of the lipid components between solid and liquid (mobile) phases. Elution of the desired lipid class is achieved by varying the polarity and strength of the mobile phase. Common stationary phases for column chromatography are silica, alumina, and ion exchange resins, whereas the preferred column materials for lipid analysis are silicic acid as well as florisil (magnesium silicate).

Low pressure column chromatography using 50- to 500-mesh adsorbents has been used commonly for separation of different lipid classes. The main parameters involved in column chromatography include weight of the adsorbent, conditioning of the adsorbent (moisture content), and column size. It is generally accepted that long narrow columns give the best resolution, but large-diameter columns increase sample capacity. For convenience, diameters over 5 cm and heights over 45 cm are not recommended for typical laboratory use [4,82].

In adsorption chromatography, compounds are bound to the solid adsorbent by polar, ionic, and, to a lesser extent, nonpolar or van der Waals forces. Therefore, separation of lipid components takes place according to the relative polarities of the individual components, which are determined by the number and type of nonpolar hydrophobic groups. In general, elution of the column with solvents with increasing polarity separates the lipid mixture according to increasing polarity of its components in the following order: saturated hydrocarbons, unsaturated hydrocarbons, wax esters, steryl esters, long chain aldehydes, triacylglycerols, long chain alcohols, FFAs, quinones, sterols, diacylglycerols, monoacylglycerols, cerebrosides, glycosyl diacylglycerols, sulfolipids, acidic glycerophosphatides, phosphatidylethanolamine, lyso-phosphatidylethanolamine, phosphatidylcholine, sphingomyelin, and lysophosphatidylcholine [83]. The procedure applicable to most lipid mixtures is eluting on a silicic acid column with the solvent sequence of chloroform, acetone, and methanol to separate into neutral lipids, glycolipids, and phospholipids, respectively [4,83].

The ion exchange columns carry ionic groups bind to the opposite charge of
the ionic groups of lipids. Thus, a polymer with fixed cations will bind anionic lipids from a mixture, provided that the pH of the solvent mixture allows ionization of the anionic groups. At the same time, the concentration of nonlipid anions in the solvent mixture should not compete for all of the fixed ions [84]. Some of the ion exchange chromatographic materials commonly used for lipid analysis are given in Table 2. Diethylaminoethyl (DEAE) cellulose is used for separation of lipid classes, and triethylaminoethyl (TEAE) cellulose is useful for separation of lipids having only ionic carboxyl groups (e.g., fatty acids, bile acids, gangliosides) or phosphatidylethanolamine from ceramide polyhexosides [83]. In polar lipid analysis, DEAE cellulose in the acetate form is the most frequently used anion exchange material. It is most effective in the pH range 3–6, and often separation of polar lipid is achieved by stepwise elution with ammonium acetate buffer in water–ethanol. The cation exchanger carboxymethylcellulose (CMC) as its sodium salt has been used occasionally over the same pH range for separation of phospholipids [84].

Complexing the adsorbent material with silver nitrate enables separation of lipid mixtures according to the number, position and cis and trans isomerism of double bonds in unsaturated fatty acids and their derivatives. Use of borate treatment of the column material complexes the compounds containing hydroxyl groups on adjacent carbon atoms and assists the separation of glycolipids [84]. Complexing of adsorbent materials is discussed in detail in the section on thin-layer chromatography.

Nowadays commercial columns are prepacked with a variety of solid stationary phases, which are available for separation of lipid classes and may be referred to as solid phase extraction (SPE) columns. SPE consumes less time, solvent, and packing material than does classical column chromatography [85]. SPE can be used for isolation, concentration, purification, and fractionation of analytes from complex mixtures [86,87]. Aminopropyl-bonded phase has been used for separation of total lipids in lipid classes obtained from different sources [88–91].

Immunoaffinity column chromatography has been used for isolation and purification of apolipoproteins. Ligand molecules (antibody; antibody to apolipoprotein or antigen; purified apolipoprotein) are immobilized on the solid support (matrix) and bind to corresponding target molecules (antigen or antibody) in a mixture of macromolecules. The bound target molecule (e.g., apolipoprotein) can then be desorbed from the ligands and eluted in the purified form using appropriate buffers [92].

<table>
<thead>
<tr>
<th>Ionizing group</th>
<th>Commercial classification</th>
<th>Analytical use</th>
</tr>
</thead>
<tbody>
<tr>
<td>$-\text{(CH}_2\text{N')H(C}_2\text{H}_5\text{)}_2$</td>
<td>DEAE (anionic exchanger)</td>
<td>Anionic lipids (phospholipids, sulfolipids, sialoglycolipids, fatty acids)</td>
</tr>
<tr>
<td>$-\text{(CH}_2\text{N')C}_2\text{H}_5$</td>
<td>TEAE (anionic exchanger)</td>
<td>Anionic lipids</td>
</tr>
<tr>
<td>$-\text{CH}_2\text{COO}^-\text{(carboxymethyl)}$</td>
<td>CM (cationic exchanger)</td>
<td>Phospholipid mixtures</td>
</tr>
</tbody>
</table>

Source: Adapted from Ref. 84.
2. Gas Chromatography

The GC (or GLC) analysis of lipids has been much studied in the literature. This method involves partitioning of the components of the lipid mixture in the vapor state between a mobile gas phase and a stationary nonvolatile liquid phase dispersed on an inert support.

Analysis of fatty acid composition by GC usually requires derivatization of fatty acids to increase their volatility. Fatty acid methyl esters (FAME) may be prepared by different transmethylation techniques and then separated on GC columns and detected by flame ionization detection (FID). The gas phase for GC is usually nitrogen or helium for packed columns and helium or hydrogen for capillary columns. The identification of chromatographic peaks is based on comparison of their retention times with those of authentic samples. GC analysis of TAG of food lipids may also provide information about positional distribution of fatty acids in the molecules. Naturally occurring TAGs that are purified by TLC can then be resolved without derivatization on the basis of their carbon number or molecular weight using capillary GC equipped with 8- to 15-m-long columns coated with methylphenyl-, methyl-, or dimethylsilicone (nonpolar capillary). Use of helium or hydrogen as a carrier gas for separation of TAG on such columns requires higher temperatures than those employed for separation of methyl esters. Mono- and diacylglycerols have to be converted to trimethylsilyl (TMS) or tert-butyldimethylsilyl ethers (TBDMS) for complete resolution [93,94]. A combined GC and mass spectrometric technique has been applied for determining molecular species in the glycerol esters. TMS or t-BDMS derivatives of glycerol esters separated on GC may be subjected to mass spectrometric analysis in order to obtain information on their molecular structure [95].

3. High-Performance Liquid Chromatography

The HPLC is a highly efficient form of adsorption, partition, or ion exchange LC that uses a very uniform, finely divided, microspherical (5–10 μm) support of controlled porosity and degree of hydration. The adsorbent is tightly packed into a stainless steel column (10–30 cm long, 2–4 mm diameter) and requires a high pressure pump to obtain an adequate and constant flow of solvent through the column. Elution of the column may be carried out either isocratically with a solvent mixture of constant composition, or by gradient elution in which the solvent composition may be varied linearly or in a stepwise fashion with both binary and ternary solvent systems. The column eluates are continuously monitored by means of a flow-through detector, which should be insensitive to solvent flow rate, temperature, and composition [83].

Sample derivatization is employed to facilitate the separation and/or to enhance the limit of detection for the HPLC analysis. Hydrolysis or saponification is done to cleave ester linkages and to obtain FFAs for their subsequent analysis. Although lipids do not possess specific UV absorption peaks, they could be detected in the region of 203–210 nm due to the presence of double bonds in the fatty acyl groups, or the functionalities such as carbonyl, carboxyl, phosphate, amino, or quaternary ammonium groups. However, this low UV range greatly restricts the choice of solvent, and it is advisable to avoid chloroform–methanol mixtures because they display a strong absorption below 245 nm. Frequently, diacylglycerols require preparation of UV-absorbing derivatives (e.g., benzoate, dinitrobenzoates, pentafluorobenzoates, and TBDMS ethers) for detection. Fatty acids can be analyzed by forming 9-anthryl-
diazomethane (ADAM) derivatives and employing a fluorescence detector. Fatty acids increase their hydrophobicity when ADAM is bound to the carboxyl group; thus, the derivatives are retained longer on a reversed phase column than fatty acids and good separation is obtained [96]. Fluorescence detectors are specific for detection and quantification of tocopherols and fluorescent derivatives of fatty acids. Evaporative light-scattering detection (ELSD) and FID have been in wide use for detection of all types of lipids following HPLC separation [93,97]. These are sometimes referred to as universal detectors. The principle of ELSD involves evaporation of mobile phase of the separated lipid fraction by a nebulizer (spray the eluent stream with a large volume of nitrogen or air) to obtain droplets of solute (lipids). These solute droplets are directed through a light source (may be a laser light source); the degree of scattering of the light is proportional to the mass of the solute [98]. The RI detection may also be used for lipid analysis [99].

Normal phase HPLC also allows the separation of normal chain and hydroxy fatty acid–containing TAG. Normal phase HPLC on silver ion–loaded anion exchange columns is currently used to resolve TAG based on their degree of unsaturation [93]. Reversed phase HPLC (using C18 columns) is also widely applied for separation and quantification of tri-, di-, and monoacylglycerols.

Since glycerol possesses a prochiral carbon, asymmetrical esterification of the primary position leads to the formation of enantiomers. Although enantiomeric TAGs cannot be resolved by normal HPLC, their diastereomeric naphthylethylurethane derivatives can be separated by HPLC on silica gel [100,101]. Use of a stationary phase with chiral moieties to separate enantiomers of mono- and diacyl sn-glycerols after derivatization with 3,5-dinitrophenylurethane (DNPU) by HPLC separation of enantiomers has been reported [102–104].

Separation of phospholipids is very laborious when TLC is used; however, HPLC provides a better means of separation and quantification. At present, use of gradient (binary or tertiary) elution on silica columns is frequently used for separation of different classes of phospholipids. Several polar solvent systems that are suitable for such separations are available [105–107]. UV detection, FID, or ELSD is suitable for phospholipid identification and quantification. Separation of glycolipids can be achieved using a silica column with a binary gradient (hexane–IPA–2.8 mM ammonium acetate) [108] or reversed phase C18 (ODS) column [98].

Isocratic, normal phase separation of cholesterol esters, FFAs, and free sterols is widely used. Simultaneous analysis of nonpolar and polar lipids using HPLC silica gel column has also been reported. In addition to normal and reversed phase methods, size exclusion HPLC has been used to separate TAG and other nonpolar lipids. This has been specifically employed for the analysis of polymerized lipids such as those generated during deep fat frying [60]. Christopoulou and Perkins [109] have described separation of monomers, dimers, and trimers of fatty acids in oxidized lipids using size exclusion column with an RI detector, whereas Burkov and Henderson [110] have reported the use of a similar column with ELSD to analyze polymers in autoxidized marine oils.

Micro-HPLC columns have the volume of one-hundredth of that of a conventional column and benefit from low consumption of material used as stationary phase, sample and mobile phase, operation at low flow rate, and temperature programming. It can also be easily coupled with a mass spectrometer and an FTIR detector [96].
4. Supercritical Fluid Chromatography (SFC)

Use of SFC for lipid extraction was discussed in a previous section. When CO₂ is compressed at a temperature and pressure above its critical point it does not liquify but forms a dense gas; thus, as a mobile phase SFC is gaseous and solvating. Such a dense gas has a number of properties (e.g., relatively high densities and diffusivities) that make it attractive for use as a mobile phase for LC. SFC with open tubular columns acts as a substitute for GC, but with the analysis temperature much lower than that employed in GC.

The temperature and pressure required for SC-CO₂ are much lower than that for HPLC. As CO₂ is nonpolar its SFC can dissolve less polar compounds and is suitable for the analysis of less polar species. To analyze polar components, polar solvents such as methanol or ethanol may be added to the SFC to cover active sites (-Si-OH) on the surface of the supporting material and to increase the dissolving power of the mobile phase. Both packed and capillary columns are used for SFC. Packing materials developed for HPLC are suitable for SFC packed columns [111]. Similarly, fused silica capillary tubes used for GC are suitable for SFC and stationary phases may employ dimethylpolysiloxane, methylphenylpolysiloxane, diphenylpolysiloxane, and cyanopropylpolysiloxane [112].

Capillary SFC with FID or UV has been used for analysis of TAG, FFAs, and their derivatives [113–115]. SFC argentation chromatography has been used for the separation of TAG according to the number of double bonds, chain length, and the nature of the double bonds [116,117]. It is necessary to add acetonitrile as a polar modifier to CO₂ to facilitate elution of TAGs. For improved analytical efficiency of lipid having a narrow range of unsaturation, mobile phase gradient (e.g., temperature, pressure and/or density, velocity) can be employed. Detectors for ELSD, FID, UV, mass spectrometry (MS), and FTIR developed for GC and HPLC are applicable to SFC. Combination of SFC with supercritical fluid extraction (SFE) has been successful for analyzing lipids of different food samples. SFE may replace any conventional lipid extraction method, and the quantification of extracted lipids (instead of gravimetric analysis) can be performed with a detector (ELSD) that has been directly connected to the extraction cell. SFE-SFC has also been used to characterize TAG patterns of seed oils [111,118]. SFC is a viable alternative for reducing any solvent use in lipid extraction and analysis and has a great potential for further development.

5. Thin-Layer Chromatography

TLC is one of the main analytical tools used for lipid analysis. TLC can be used for fractionation of complex lipid mixtures, assay of purity, identification, information on the structure, as well as for monitoring extraction and separation of components via preparative column chromatography for routine and experimental purposes. The principles and theory of TLC are based on the difference in the affinity of a component toward a stationary and a mobile phase. The important components of TLC are the stationary phase, mobile phase, detection, and quantification [5]. The adsorbent generally used in TLC for lipid analysis is very fine-grade silica gel and may contain calcium sulfate as a binder to ensure adhesion to the plate [4]. Alumina and kieselguhr are also used as stationary phases. These adsorbents can also be modified by impregnation with other substances so as to achieve the desired separations. The most popular impregnations are with boric acid or silver nitrate. Silver nitrate—
impregnated (argentation) TLC may be used to separate TAGs or fatty acid methyl esters according to the number of double bonds and also by virtue of the geometry (e.g., cis or trans) and position of the double bonds in the alkyl chain. The silver ion forms a reversible complex with the π electrons of the double bond of unsaturated fatty acids, thereby decreasing their mobility [119]. Structural isomers of TAGs (due to their fatty acid constituents) may also be separated on this type of TLC plate [97]. Impregnation of the TLC plates with boric acid (3%, w/v) prevents isomerization of mono- and diacylglycerols while separating neutral lipids [83]. Boric acid complexes with vicinal hydroxyl groups and leads to slower migration of these compounds [119].

Samples of lipid extracts are applied as discrete spots or as narrow streaks, 1.5–2 cm from the bottom of the plate. The plate is then developed in a chamber containing the developing solvent or a solvent mixture. The solvent moves up the plate by capillary action, taking the various components with it at different rates, depending on their polarity and how tightly they might be held by the adsorbent. The plate is removed from the developing chamber when the solvent approaches the top of it and then dried in the air or under a flow of nitrogen. Solvents with low boiling point, viscosity, and toxicity are suitable for TLC application. A low boiling point helps in the quick evaporation of the solvent from the surface layer, and low viscosity facilitates faster movement of the solvent during development. The selection of a suitable solvent is very important for good separation of the lipid classes. Several solvent systems may be used to resolve individual lipid classes, as exemplified in Table 3.

The location of the corresponding lipid spots on a developed plate has to be detected prior to their isolation. Detection of the spots may be done using a reagent directly on the plate. This reagent could be specific to certain functional groups of the lipid molecules or may be a nonspecific reagent that renders all lipids visible. There are nondestructive chemical reagents, such as 2′,7′-dichlorofluorescin in 95% methanol (1%, w/v), iodine, rhodamine 6G, and water, which allow recovery of lipids after detection. Lipids exhibit a yellow color and in the presence rhodamine 6G (0.01%, w/v) produce pink spots under UV light. These chemicals may also be used as a nondestructive spray for preparative TLC. When water is used separated lipids may appear as white spots in a translucent background and can easily be distinguished. Developed plates may also be subjected to saturated iodine vapor in a chamber and this may produce brown spots due to the reaction of iodine with unsaturated bonds of the lipid molecules. However, unsaturated lipids may form artifacts with iodine, if sufficient time is allowed. The destructive methods include spraying of the plate with sulfuric acid (50%, v/v) and drying at 180°C for 1 hour to make lipids visible as black deposits of carbon [4]. Potassium dichromate (5%, w/v) in 40% (v/v) sulfuric acid also works in a similar manner to sulfuric acid spray. Molybdophosphoric acid (5%, w/v) in ethanol turns lipids into blue then black when heated at 120°C for 1 hour. Coomassie blue (0.03% in 20% methanol) turns lipid into blue spots on white background. Examples of specific reagents that react selectively with specific functional groups include FeCl₃ to detect cholesterol and cholesterol esters, ninhydrin for choline-containing phospholipids, and orcinol or naphthol/H₂SO₄ for glycolipids [123]. Some lipids contain chromophores and can be visualized directly under UV or visible light without staining.
<table>
<thead>
<tr>
<th>Lipid component</th>
<th>TLC adsorbent</th>
<th>Solvent system</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complex lipids (animal tissues)</td>
<td>Silica gel G</td>
<td>Chloroform–methanol–water, 25:10:1, v/v/v</td>
<td>120</td>
</tr>
<tr>
<td></td>
<td>Silica gel H</td>
<td>Chloroform–methanol–acetic acid–water, 25:15:4:2, v/v/v/v</td>
<td>120</td>
</tr>
<tr>
<td></td>
<td>Silica gel H</td>
<td>First developing system, pyridine–hexane, 3:1, v/v and second developing system, chloroform–methanol–pyridine–2 M ammonia, 35:12:65:1, v/v/v/v</td>
<td>120</td>
</tr>
<tr>
<td>Complex lipids (plant tissues)</td>
<td>Silica gel G</td>
<td>Acetone–acetic acid–water, 100:2:1, v/v/v</td>
<td>121</td>
</tr>
<tr>
<td>Simple lipids</td>
<td>Silica gel G</td>
<td>Diisobutyl ketone–acetic acid, 40:25:3:7, v/v/v</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Silica gel G</td>
<td>Hexane–diethyl ether–formic acid, 80:20:2, v/v/v</td>
<td>4, 122</td>
</tr>
<tr>
<td></td>
<td>Silica gel G</td>
<td>Benzene–diethyl ether–ethyl acetate–acetic acid, 80:10:10:2, v/v/v/v</td>
<td>4, 122</td>
</tr>
<tr>
<td>Partial acylglycerol</td>
<td>Silica gel G containing 5% (w/v) boric acid</td>
<td>Chloroform–acetone, 96:4, v/v</td>
<td>4</td>
</tr>
<tr>
<td>Neutral plasmalogens</td>
<td>Silica gel G</td>
<td>Hexane–diethyl ether, 95:5, v/v, in first direction, hexane–diethyl ether, 80:20, v/v, in second direction</td>
<td>4</td>
</tr>
</tbody>
</table>

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Lipids detected by nondestructive methods can be recovered by scraping the bands of interest and dissolving them in an appropriate solvent. Complex lipid mixtures cannot always be separated by one-dimensional TLC; however, two-dimensional TLC may resolve such mixtures.

The separation efficiency of TLC plates is affected by the degree of hydration of the adsorbent. Therefore, activation of the adsorbent, which depends on both the time and the temperature as well as the storage conditions of the plates and the relative humidity of the atmosphere, must be considered. Use of authentic lipid standards would allow direct comparison of the resolved lipids in an unknown mixture. High-performance TLC (HP-TLC) plates have high resolving power and speed of separation. They are available commercially as precoated plates with fine (5-μm) and uniform particle size silica gel. However, the amount of sample that can be applied to such HP-TLC plates is very small. TLC is a preferred method of purification and separation of lipid classes before subjecting them to further separation of individual components.

For quantification of TLC separated lipids, traditional scraping followed by quantitation and in situ determination are available. The separated lipid classes on silica gel can be scraped off, extracted by means of suitable solvents, and quantitated gravimetrically, spectrophotometrically, or by GC. Determination of the phosphorus content of the eluted phospholipids is a classic example for spectrophotometric quantitation. GC can be employed to quantify separated neutral and phospholipid classes by derivatizing the constituent fatty acids into their methyl esters. Densitometric methods provide an in situ quantification method for lipids. Lipids are sprayed with reagents and their absorption or fluorescence can be measured under UV or visible light by densitometry.

The spots detected by charring can be measured by scanning photodensitometer areas of peaks, which are proportional to the amount of original lipid present, are recorded. Scintillation counting is also possible after introducing a correct scintillator (e.g., mixture of 2,5-diphenyloxazole and 1,4-bis-2-(5-phenyloxazoyl)benzene, PPO, and POPOP in toluene) into the lipid. Scanning fluorometry allows resolution of lipids on an adsorbent containing a fluorescent dye or by spraying with such reagents.

TLC could be coupled with other methods to facilitate detection, quantitative identification, or quantitation of separated lipids. These include coupling with HPLC (HPLC/TLC), Fourier transform infrared (TLC/FTIR), nuclear magnetic resonance (TLC/NMR), and Raman spectroscopy (TLC/RS) [119]. The Iatroscan (Iatron Laboratories of Japan) applies the same principles of TLC to separate lipid mixtures followed by their detection using FID. The TLC medium is silica (7.5 μm thick) sintered onto quartz rods (0.9 mm in ID, 15 cm long, called Chromarods). Chromatographic resolution of lipid classes on Chromarods and the composition of solvent systems used are similar to those employed in classical TLC with modifications. Copper sulfate, silver nitrate, boric acid, or oxalic acid impregnation has been reported to produce better resolution and increased responses in the determination [124]. Parrish and Ackman [125] have shown that stepwise scanning and developing in solvent systems of increasing polarity resolves individual lipid components in neutral lipids of marine origin. FID gives a linear response that is proportional to the number of nonoxidized carbon atoms in the material entering the flame. The Chromarods are reusable up to 100–150 times and require thorough cleaning after each use. The main advantage of TLC-FID is the short analysis time, with the pos-
UV and visible spectroscopy are less frequently used but have specific applications for the identification and quantification of lipids. IR spectroscopy was the first spectroscopic method applied for the analysis of lipids. NMR and MS have been widely used for lipid structure determination; however, new applications other than these have been developed. IR, UV, and NMR are nondestructive spectroscopic methodologies.

1. **UV-Visible Spectroscopy**

UV and visible spectra of organic compounds are attributable to electronic excitations or transitions. Functional groups with high electron density, such as carbonyl and nitro groups with double, triple, or conjugated double bonds, absorb strongly in the ultraviolet or visible range at characteristic wavelengths ($\lambda_{\text{max}}$) and molar extinction coefficients ($\varepsilon_{\text{max}}$). Table 4 provides some of the diagnostic UV absorption bands for lipid analysis. It should be noted that the $\lambda_{\text{max}}$ for a compound may vary, depending on the solvent used [83,84].

2. **Infrared Absorption Spectroscopy**

The IR spectrum of an oil provides substantial information on the structure and functional groups of the lipid and also about the impurities associated with it. These information are represented as peaks or shoulders of the spectrum as illustrated in Figure 1. At the high frequency end of the spectrum (3700–3400 cm$^{-1}$), there is a

### Table 4  UV Absorption Characteristics of Some Chromophoric Groups

<table>
<thead>
<tr>
<th>Chromophore</th>
<th>Example</th>
<th>$\lambda_{\text{max}}$ (nm)</th>
<th>$\varepsilon_{\text{max}}$</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>$-\text{C}=-\text{C}-$</td>
<td>Octene</td>
<td>177</td>
<td>12,600</td>
<td>Heptane</td>
</tr>
<tr>
<td>$-\text{C}/\text{C}-$</td>
<td>Octyne</td>
<td>178</td>
<td>10,000</td>
<td>Heptane</td>
</tr>
<tr>
<td>$-\text{C}=-\text{C}=-\text{C}-$</td>
<td>Butadiene</td>
<td>196</td>
<td>2,100</td>
<td>Heptane</td>
</tr>
<tr>
<td>$-(\text{C}=-\text{C})_n-$</td>
<td>Conjugated polyenes</td>
<td>$217 + 30 \ (n - 2)$</td>
<td>20,000–100,000</td>
<td>Hexane</td>
</tr>
<tr>
<td>C$_6$H$_6$</td>
<td>Benzene</td>
<td>184</td>
<td>47,000</td>
<td>Cyclohexane</td>
</tr>
<tr>
<td>C$_6$H$_5$</td>
<td>$\beta$-Carotene</td>
<td>452</td>
<td>139,000</td>
<td>Hexane</td>
</tr>
<tr>
<td>HC$\equiv$O</td>
<td>Acetaldehyde</td>
<td>290</td>
<td>17</td>
<td>Hexane</td>
</tr>
<tr>
<td>C$\equiv$O</td>
<td>Acetone</td>
<td>275</td>
<td>17</td>
<td>Ethanol</td>
</tr>
<tr>
<td>$\text{C}=\text{O}$</td>
<td>Acetic to palmitic acid</td>
<td>208–210</td>
<td>32–50</td>
<td>Ethanol</td>
</tr>
</tbody>
</table>

Adapted from Ref. 83.
Figure 1  The mid-infrared spectrum of an edible oil collected on an attenuated total reflectance crystal. Labels indicate absorption bands or regions associated with triacylglycerols or other constituents that may be present in an oil or free fatty acid. (From Ref. 137.)

region where compounds containing hydroxyl groups are absorbed, including water (H-OH) and hydroperoxides (ROH-H) and their breakdown products. At some lower frequencies (3025–2850 cm\(^{-1}\)) is the CH stretching region where three bands are visible: a weak cis double bond CH absorption (CH\(\equiv\)CH) and strong bonds due to the CH\(_2\) groups of the aliphatic chains of TAGs as well as the terminal methyl groups. Just beyond this region is secondary oxidation products of lipids such as aldehydes and ketones that absorb energy albeit weakly. Toward the center of the spectrum is a very strong band due to the C=O stretching absorption due to the ester linkage attaching the fatty acids to the glycerol backbone of TAGs. Next to it is a band due to COOH group of FFAs if the lipid is hydrolyzed. In the same region there would be carbonyl absorption bands of aldehydes (R-CHO) and ketones (R-CO-R) if the oil is oxidized. Continuing to lower frequencies area is the finger printing region (1500–900 cm\(^{-1}\)) as the pattern of bands in this region is very characteristic of molecular composition of the lipid and could be used to identify different components. At the low-frequency end of the fingerprinting region, a band due to the CH\(\equiv\)CH bonding absorption of isolated trans double bonds could be seen when trans-containing TAGs are present in the oil. The corresponding absorption of conjugated dienes containing trans-trans and cis-trans double bonds appears at slightly.
the isolated trans band is another group of CH absorption bending vibrations including a very strong cis absorption [137].

The IR absorption signal could be employed to analyze and obtain information about qualitative structural and functional groups of lipids. In principle, since IR band intensities are linearly related to the concentration of the absorbing molecular species, quantitative information about the lipid can also be obtained [137]. IR spectroscopy has been applied to solid lipids to obtain information about polymorphism, crystal structure, conformation, and chain length. In oils, IR is commonly used to determine the presence and the content of trans unsaturation. Single trans double bonds show a characteristic absorption band at 968 cm\(^{-1}\), and the frequency does not change for additional double bonds unless these are conjugated. There is no similar diagnostic IR absorption band for cis unsaturation; however, Raman spectra show strong absorption bands at 1665 ± 1 cm\(^{-1}\) (cis-olefin), 1670 ± 1 cm\(^{-1}\) (trans-olefin) and 2230 ± 1 and 2291 ± 2 cm\(^{-1}\) (acetylene) for the type of unsaturation shown [94]. Kates [83] has provided the characteristic IR absorption frequencies that have diagnostic values for identification of major classes of lipids. It has also been reported that ionic forms of phospholipids influence the absorption bands associated with phosphate groups that influence the interpretation of the spectra [83]. The FTIR spectrometer finds its uses in measuring iodine value, saponification value, and FFAs [138]. Oxidative stability of lipids as reflected in the formation of peroxides and secondary oxidation products may also be determined by FTIR [137,139].

### 3. Nuclear Magnetic Resonance Spectroscopy

Low-resolution pulsed \(^1\)H NMR spectroscopy is employed to determine solid-fat content of lipids as well as the oil content of seeds as discussed earlier in this chapter. High-resolution \(^1\)H NMR applied to vegetable oils gives several signals with designated chemical shifts, coupling constant, splitting pattern, and area. This information can be used to obtain structural and quantitative information about lipids. Methyl stearate (saturated fatty acid ester) may give five distinct \(^1\)H NMR signals as summarized in Table 5. Similar signals appear in methyl oleate and linoleate, but methyl oleate also gives signals for olefinic (5.35 ppm, 2H) and allylic (2.05 ppm, 4H) hydrogen atoms, and for linoleate these are at 5.35 (4H), 2.05 (4H, C8, and C14) and 2.77 ppm (2H, C11). When a double bond gets close to the methyl group, as in α-linolenate and other ω-3 esters, the CH\(_3\) signal is shifted to 0.98 ppm; oils

### Table 5  Chemical Shift (ΔH) for Methyl Alkanoates Observed for \(^1\)H NMR

<table>
<thead>
<tr>
<th>ΔH (ppm)</th>
<th>Splitting</th>
<th>H</th>
<th>Group$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.90</td>
<td>Triplet</td>
<td>3</td>
<td>CH(_3)</td>
</tr>
<tr>
<td>1.31</td>
<td>(Broad)</td>
<td>2n</td>
<td>(CH(_2))(_n)</td>
</tr>
<tr>
<td>1.58</td>
<td>Quintet</td>
<td>2</td>
<td>—CH(_2)CH(_2)COOCH(_3)</td>
</tr>
<tr>
<td>2.30</td>
<td>Triplet</td>
<td>2</td>
<td>—CH(_2)CH(_2)COOCH(_3)</td>
</tr>
<tr>
<td>3.65</td>
<td>Singlet</td>
<td>3</td>
<td>—CH(_2)CH(_2)COOCH(_3)</td>
</tr>
</tbody>
</table>

$^a$Assigned hydrogen is designated as H.

Adapted from Ref 94
containing other esters (ω-6, ω-9, and saturated) exhibit a triplet at 0.89 ppm. The area associated with these various signals can be used to obtain semiquantitative information in terms of ω3 fatty acids (α-linolenate), other polyenic, monoenic (oleate), and saturated acids [94,140].

Acylglycerols show signals associated with the five hydrogen atoms on the glycerol moiety. There is a one-proton signal at 5.25 ppm (CHOCOR), which overlaps with the olefinic signals and a four proton signals located between 4.12 and 4.28 ppm (CH₂OCOR). Phospholipids display characteristic signals for phosphatidylcholine and phosphatidylethanolamine [94].

High-resolution ¹³C NMR spectra are more complex than ¹H spectra and provide more structural information rather than quantitative ones. It is also possible to locate functional groups such as hydroxy, epoxy, acetylenic, and branched chains in the molecules. The application of ¹³C NMR to TAG allows determination of the positional distribution of fatty acids on the glycerol backbone [141,142]. The ¹³C resonance of the carbonyl group of fatty acids in the sn-1 and sn-3 positions is well resolved from those esterified at the sn-2 position. Most unsaturated fatty acids in the sn-2 position are nondegenerate and could be easily differentiated. ¹³C NMR has been successfully applied for determination of positional distribution of TAG fatty acids in vegetable oils [143–145] and marine oils [146,147].

The NMR imaging is based on manipulation of magnetic field gradients oriented at right angles to each other to provide spatial encoding of signals from an object, which are converted by FT techniques to three-dimensional NMR images [148]. It produces three-dimensional data by selecting two-dimensional cross-sections in all directions. Application of NMR imaging or magnetic resonance imaging (MRI) to foods has been of interest as it is a noninvasive technique that can be applied to track the dynamic changes in foods during storage, processing, packaging, and distribution.

Most magnetic resonance images of foods are based on proton resonances from either water or lipids. Simoneau et al. [149] have applied MRI to the study of fat crystallization in bulk or dispersed systems. Halloin et al. [150] described two MRI techniques, spin-echo imaging (SEI) and chemical shift imaging (CSI), for the study of lipid distribution in pecan embryos. Insect- or fungus-damaged embryos gave images that were less intense than those of normal embryos, reflecting lower oil content. When MRI and NMR was employed to determine the oil content of French-style salad dressings, results were within ±2% of expected values and were in agreement with oil content determined by traditional methods [151]. Pilhofer et al. [152] have studied the use of MRI to investigate the formation and stability of oil-in-water emulsions formed with vegetable oil, milk fat, and milk fat fractions. Distribution of lean and fat in retail meat as a means of quality can be measured [153] using MRI and also to visualize oil and water concentration gradient during deep-fat frying food [154].

4. Mass Spectrometry

In conventional MS, compounds in their gaseous state are ionized by bombardment with electrons (electron impact) in an ionization chamber. The resulting mass spectrum consists of a characteristic pattern of peaks representing molecular fragments with different mass to charge (m/z) ratios. Some of these peaks or patterns of peaks are structurally diagnostic. The parent ion peak that arises from the unfragmented
ionized molecule has the highest m/z ratio, but may not be always present, depending on the volatility and thermal stability of the compound. Thus, lipids containing polar groups, such as phospholipids, with low thermal stability and volatility and high molecular weight cannot be analyzed by conventional electron impact (EI) mass spectrometry. Therefore, fast atom bombardment (FAB), chemical ionization (CI), field desorption (FD), or secondary ion (SI) mass spectrometry is required for such lipid analysis [95].

MS is very useful in identifying structural modification of chain length such as branching or the presence of rings for saturated species. In this regard, matrix-assisted laser desorption ionization and time-of-flight mass spectrometry (MALDI-TOF-MS) have several advantages. It does not require prior derivatization of sample to enhance the volatility of the lipids. The extent of fragmentation of MALDI-TOF-MS is low; thus, detection of molecular ion is possible in most cases. Schiller et al. [155] have successfully used a matrix of 2,5-dihydroxybenzoic acid to identify phosphatidylcholine and different phospholipids as their molecular ions (M + 1). Diacylglycerols were mainly detected as their corresponding sodium or potassium adducts but not as their protonated form. MALDI-TOF-MS can be used for direct investigation of lipid mixtures occurring, e.g., in cell membranes due to its high sensitivity (up to picomolar concentrations).

MS in combination with GC and HPLC is also useful in structural determination of the individual lipid molecules. Le Que´re´ et al. [156] have developed an on-line hydrogenation method that allows selective hydrogenation of all the unsaturated species after chromatographic separation for deducing structural information such as carbon skeleton and double-bond equivalents. Le Que´re´ [157] has reviewed this methodology and the use of GC-MS and tandem MS for analysis of structural features of fatty acids.

D. Enzymatic Methods

Higgins [158] has described an enzymatic method for determining TAG content of samples. This involves reaction of the TAG with lipase in order to obtain glycerol and FFAs. The glycerol so produced is then converted to α-glycerophosphate using glycerol kinase. The α-glycerophosphate dehydrogenase is then used to reduce nicotinamide adenine diphosphate (NAD) to NADH. The resultant NADH is then measured by a colorimetric reaction.

Cholesterol oxidase is used for determining cholesterol concentration in blood plasma. Polysaturated fatty acids (PUFA) with cis-methylene groups between their double bonds (e.g., linoleic, linolenic, and arachidonic acids) can be quantitatively measured by reading the UV absorbance of conjugated diene hydroperoxides produced via lipoxygenase (lipoxidase)–catalyzed oxidation. The extinction coefficient of diene hydroperoxides at 234 nm is the same for all PUFAs. Fatty acid esters have to be saponified prior to such analysis. The phosphatidylcholine or lecithin content of foods (e.g., as a measure of the egg content of foods) can be made by catalyzing conversion of lecithin to phosphatidic acid and choline by lecithinase (phospholipase D) [5].

The method of stereospecific analysis of TAG described by Brockerhoff and Yurkowski [159] uses pancreatic lipase that eventually removes fatty acids from the sn-1 and sn-3 positions of the TAG. This procedure has recently been employed to
determine the existing structural differences of fish oils and seal oil [160]. Phospho-
lipase A₂ is used to release fatty acids at the sn-2 position of the synthesized phos-
phatide during the analytical procedure. Lands [161] used a different approach to
determine steric positions of the fatty acids. TAG is hydrolyzed with lipase and the
products are separated by TLC. The sn-3 hydroxyl of glycerol is phosphorylated with
diacylglycerol kinase to produce 3-phosphoryl monoacylglycerol. In the following
step of analysis, phospholipase A₂ is used to remove the fatty acids only from the
sn-2 position. The fatty acids in the sn-1 position can be released by saponifying the
resultant 1-acyl lysophosphatidic acid. The fatty acids in positions 1 and 2 can be
identified by GC analysis. The composition of fatty acids in position 3 can be cal-
culated by comparing these results with those from total fatty acid composition
determination of TAG.

E. Immunochemical Methods

Lipids are not generally very immunogenic. However, most glycolipids (except in
the pure form) possess antibodies of high activity and specificity. Therefore, glyco-
lipids to be administered to the animal are conjugated by covalent linking to a foreign
protein (as a hapten) or using them as a part of the bilayer of a liposome to stimulate
the production of specific antibodies [55,57]. Immunochemical methods have also
been developed for the assay of phospholipids and TAGs (55). The steroid hormones
when conjugated with serum albumin are sufficiently immunogenic to stimulate gen-
eration of antibodies with high activity, and this allows their detection.

Immunostaining of TLC plates for detection and assay of glycoproteins is
widely done. The TLC chromatogram containing separated glycolipids is treated with
a radiolabeled specific antibody (usually with ¹²⁵I) to stain only the glycolipid antigen
even in the presence of overlapping glycolipids. Detection of ¹²⁵I may be achieved
using autoradiography, and the chromatographic mobility and antibody staining
serves to identify the glycolipid [84,92]. To overcome low sensitivity of the immu-
noradiolabeled detection of glycolipids, enzyme-linked immunosorbent assay
(ELISA) was developed. For ELISA, lipid is usually bound to a solid phase and the
antibody is measured either by virtue of itself carrying enzyme or by using a second
antibody that carries an enzyme [84].

V. SUMMARY

Lipids are integral components and building blocks of biological materials. To un-
derstand their constituents, chemistry, and biological functions, lipids have to be
isolated and studied. Therefore, an extensive knowledge of the extraction and anal-
ysis of lipids is essential to carry out studies on lipids. This chapter provided com-
prehensive information on methods available for extraction and analysis of lipids
from biological materials with examples when necessary. More details of a particular
topic could be obtained from the references listed.

REFERENCES

   for Nutrition Labeling* (D. M. Sullivan and D. E. Carpenter, eds.). AOAC Press, Ar-
  lington, VA, 1993, pp. 85–104.


Methods for \textit{trans} Fatty Acid Analysis

RICHARD E. MCDONALD
U.S. Food and Drug Administration, Summit-Argo, Illinois

MAGDI M. MOSSOBA
U.S. Food and Drug Administration, Washington, D.C.

I. INTRODUCTION

\textit{trans} Fatty acids are present in a variety of food products; some are derived from natural sources, such as dairy products, but most come from products that contain commercially hydrogenated fats. The nutritional properties of \textit{trans} fatty acids have been debated for many years, particularly with respect to the amounts of low density and high density lipoprotein (LDL, HDL) contained in serum. Some studies have shown that \textit{trans} fatty acids elevate levels of serum LDL cholesterol and lower HDL cholesterol [1–3]. Such results drew a great deal of media attention, which led to several requests for the U.S. Food and Drug Administration (FDA) to make labeling of \textit{trans} fatty acids mandatory on food products. There have also been requests to either ban these substances or to impose strict limitations on their use. However, a comprehensive report of an expert panel convened by the International Life Sciences Institute [4] concluded that food products containing partially hydrogenated fats are good substitutes for traditional fats rich in saturated fat, but are not good substitutes for unhydrogenated vegetable oils. The report also concluded that the overall effect of hydrogenated fat on serum cholesterol levels depends on the consumption of saturated, \textit{trans}/\textit{cis}-monounsaturated, and \textit{cis}-polyunsaturated fatty acids. The authors recommended more research to better understand the effects of \textit{trans} fatty acids on serum lipid concentrations and coronary heart disease. We recently discussed both the nutritional and food labeling issues associated with \textit{trans} fatty acids [5].

\textit{trans} Fatty acids in the diet are mainly derived from partially hydrogenated vegetable oils (PHVOs) and ruminant fats. While PHVOs have been reported to
present a possible risk factor for coronary heart disease [6], some trans conjugated fatty acids in ruminant fat have been reported to have several beneficial physiological effects in experimental animals [7]. trans-Vaccenic acid (trans-11-18:1), which is the major trans fatty acid isomer present in meat and dairy products from ruminants [8,9], has been shown to be converted to cis-9,trans-11-18:2, a conjugated linoleic acid (CLA) isomer [10–12], by the action of Δ9 desaturase present in mammalian tissue [13,14]. The relationship of trans-18:1 fatty acid isomers and CLA isomers is an active area of research today [15].

Even after several years of research, there are still several laboratories in the world that have active research programs trying to improve the methodology to determine trans fatty acids in food products. The quantitation and identification of trans fatty acid isomers is difficult because of the wide range of positional monoene, diene, and triene fatty acid isomers present in hydrogenated oils. Moreover, cis positional isomers are also present, and commercial chromatographic standards are lacking for many fatty acid isomers.

Analytical procedures used to quantify and identify fatty acids have been reviewed [5,16,17]. This chapter discusses the newest developments in analytical methods, including infrared (IR), Raman, and nuclear magnetic resonance (NMR) spectroscopy, gas chromatography (GC), high-performance liquid chromatography (HPLC), supercritical fluid chromatography (SFC), and mass spectrometry (MS). The analysis of trans fatty acid isomers is extremely challenging and complex. Thus various combinations of techniques have effectively been used to determine the concentration and confirm the identity of individual trans and cis isomers. Improved methods are still needed to accurately and conveniently determine total trans content as well as specific fatty acid isomers in hydrogenated oils.

II. ANALYSIS USING SPECTROSCOPY

A. Infrared Spectroscopy

Infrared spectroscopy is a widely used technique for determining nonconjugated trans unsaturation in both natural and processed fats and oils. It is not applicable to materials that have functional groups with absorption bands close to 966 cm⁻¹, which is the strong absorption band arising from the C—H deformation about a trans double bond. This absorption band is absent in natural vegetable oils that are composed of saturated fatty acids and fatty acids with only cis-unsaturated double bonds. For increased accuracy, oil samples are usually converted to methyl esters prior to analysis. This eliminates interfering absorptions associated with the carboxyl groups of free fatty acids and the glycerol backbone of triacylglycerols.

1. Conventional IR Methods

The early Official Method of the American Oil Chemists’ Society (AOCS), Cd 14-61 [18] for the determination of trans fatty acid concentrations in fats and oils, was based on a comparison of absorption at 966 cm⁻¹ of standards and unknowns. Samples and standards were diluted in carbon disulfide and placed in an absorption cell so that the transmittance or absorbance could be measured in an infrared spectrophotometer. The quantitation of the trans concentration was based on Beer’s law:
\[ A = abc \]  \hspace{1cm} (1)

where

\[ A = \text{absorbance} = \log \frac{1}{\text{transmittance}} \]

\[ a = \text{absorptivity} \]
\[ b = \text{path length} \]
\[ c = \text{trans concentration} \]

Disadvantages of using this method include (1) the need to make methyl ester derivatives at trans levels less than 15%, (2) the use of the toxic, volatile solvent carbon disulfide, (3) the high bias found for triacylglycerols, and (4) low accuracy obtained for trans levels less than 5%. Therefore, there has been a great deal of interest in improving methods to determine trans fatty acid concentrations.

A study was conducted that showed that the AOCS method produced trans values that were 2–3% too high, while derivatized methyl esters produced values that were 1.5–3% too low [19]. In 1965 correction factors were suggested that were incorporated into AOAC method 965.35 [20]. The percentage of trans was calculated with a procedure similar to AOCS method Cd 14-61. Several formulas using correction factors were proposed to calculate trans concentration as methyl esters or triacylglycerols. Different formulas were also used for oils containing long chain or short/medium chain fatty acids.

In a 1969 study [21], the concentration of trans fatty acids was determined based on the ratio of the IR absorptions at 965 and 1170 cm\(^{-1}\). In 1971 Firestone and Huang [22] proposed a dual-beam differential spectrophotometry procedure with a zero trans-containing vegetable oil in the reference cell. This procedure, which resulted in a 1–2% high bias for both triacylglycerols and methyl esters, was the first successful attempt to eliminate the sloping background of the trans infrared band.

In 1982 a two-component calibration curve was proposed [23] in an attempt to overcome some of the drawbacks of the earlier procedures. A calibration curve was developed by means of different levels of the trans monoene, methyl elaidate, and methyl linoleate dissolved in carbon disulfide. The calibration standards and the test samples in carbon disulfide were scanned against a carbon disulfide background and recorded from 900 to 1500 cm\(^{-1}\). After a baseline had been drawn as a tangent from about 935 and 1020 cm\(^{-1}\) at the peak minima, the corrected absorbance of the calibration standards trans peaks, at 966 cm\(^{-1}\), was obtained. The baselines for the test samples spectra were obtained by overlaying the spectra of the calibration standards spectra at corresponding concentrations. This method compensates for the low bias of other methods and eliminates the need for correction factors.

An absorption band–height ratioing procedure was used with an attenuated total reflection (ATR) cell, which allowed the use of neat samples and eliminated the need to rely on volatile toxic solvents [24]. The introduction of Fourier transform infrared spectroscopy (FTIR) instruments also facilitated more accurate and rapid determination of trans fatty acids. A laser light source was used for wavelength accuracy. Interferometers were used that allowed all wavelengths of light to be measured simultaneously. Since FTIR spectrometers are computerized, multiple spectral scans could be averaged in a few seconds. An FTIR spectrometer equipped with a
thin (0.1-mm) transmission flow cell was used to develop an automated procedure for calculating the percentage of \textit{trans} in fats and oils [25].

Postmeasurement spectral subtraction manipulations were used to correct for the highly sloping background in the FTIR absorption spectrum of hydrogenated vegetable oil fatty acid methyl esters [26]. Determination of \textit{trans} content, however, required additional IR measurement of an appropriate reference material and the digital subtraction of this reference absorption spectrum from that of the test portion. FTIR spectroscopy, in conjunction with a transmission flow cell, was used to rapidly determine the \textit{cis} and \textit{trans} content of hydrogenated oil simultaneously [27].

The 1995 official method of the AOCS for determination of the \textit{trans} fatty acid Cd 14-95 [28] utilizes two standard curves [23]. The choice of curve depends on the \textit{trans} concentration. This method is reported to be accurate to determine the \textit{trans} content of fats with \textit{trans} levels of 0.5\% or greater. Test samples and standards are converted to methyl esters, diluted in 10 mL of carbon disulfide, and then placed in a transmission IR cell before measuring the transmittance or absorbance over the range 1050–900 cm\(^{-1}\) in an infrared spectrophotometer. A typical infrared absorption spectra for an oil containing both 2\% and 70\% \textit{trans} is shown in Figure 1. The baseline-corrected absorbance (\(A_c\)) is determined by subtracting the absorbance of the baseline at the peak maximum (\(A_p\)) from the maximum absorbance at the peak (\(A_0\)). Two plots are then constructed by using standards to cover samples with low levels of \textit{trans} (1–10\%) and moderate to high \textit{trans} levels (10–70\%), and two regression equations are generated. The percentage of \textit{trans} as methyl elaidate of the unknown is then obtained by referring to the appropriate calibration data (\(\leqslant 10\%\) or >10\%) and solving the following equations:

\[
A_c = (A_p - A_b) \quad (2)
\]

Methyl elaidate weight equivalents (g) = \(\frac{A_c - \text{intercept}}{\text{slope}}\) \quad (3)

\textbf{Figure 1}  Infrared absorption spectra of fatty acid methyl esters containing 2\% and 70\% \textit{trans}. (From Ref. 28.)
\[
\%\ trans = \frac{\text{methyl elaidate weight equivalents (g)}}{\text{sample weight (g/10 mL of CS\textsubscript{2})}} \times 100
\]  

The major advantage of this method is its accuracy at low \textit{trans} levels. However, methyl ester derivatization and the use of carbon disulfide are still required.

More accurate results than those produced by the current official methods are claimed for an IR procedure, which uses a partially hydrogenated vegetable oil methyl ester mixture as the calibration standard [29]. The improved results were attributed to the assortment of \textit{trans} monoene and polyene isomers in the calibration standard with different absorbivities relative to that of methyl elaidate.

2. “Ratioing” of Single-Beam FTIR Spectra

Figure 1 indicates that the 966 cm\textsuperscript{-1} \textit{trans} band is only a shoulder at low levels. This is due to the overlap of the \textit{trans} band with other broad bands in the spectrum, which produces a highly sloped background that diminishes the accuracy of the \textit{trans} analysis. Many reports in the literature have proposed changes to the procedures above, including minor refinements to major modifications aimed at overcoming some of the limitations already discussed. These studies have resulted in the development of procedures that use spectral subtraction to increase accuracy, as well as means of analyzing neat samples to eliminate the use of solvents.

Mossoba et al. [30] recently described a rapid IR method that uses a Fourier transform infrared spectrometer equipped with an attenuated total reflection cell for quantitating \textit{trans} levels in neat fats and oils. This procedure measured the 966 cm\textsuperscript{-1} \textit{trans} band as a symmetric feature on a horizontal background. The ATR cell was incorporated into the design to eliminate one potential source of error: the weighing of test portions and their quantitative dilution with the volatile CS\textsubscript{2} solvent. The high bias previously found for triacylglycerols has been attributed to the overlap of the \textit{trans} infrared band at 966 cm\textsuperscript{-1} with ester group absorption bands. Errors for the determination of \textit{trans} concentrations below 5% that resulted from this overlap could result in relative standard deviation values greater than 50% [31]. The interfering absorption bands were eliminated, and baseline-resolved \textit{trans} absorption bands at 966 cm\textsuperscript{-1} were obtained by “ratioing” the FTIR single-beam spectrum of the oil or fat being analyzed against the single-beam spectrum of a reference material (triolein, a mixture of saturated and \textit{cis}-unsaturated triacylglycerols or the corresponding unhydrogenated oil). This approach was also applied to methyl esters. Ideally, the reference material should be \textit{trans}-free oil that has an otherwise similar composition to the test sample being analyzed.

The simplified method just outlined allowed the analysis to be carried out on neat analytes that are applied directly to the ATR crystal with little or no sample preparation. With this method, the interference of the ester absorptions with the 966 cm\textsuperscript{-1} \textit{trans} band and the uncertainty associated with the location of the baseline were eliminated. Figure 2 shows the symmetric spectra that were obtained when different concentrations of methyl elaidate (ME) in methyl oleate (MO) were “ratioed” against methyl oleate. A horizontal baseline was observed, and the 966 cm\textsuperscript{-1} band height and area could be readily measured. The minimum identifiable \textit{trans} level was 0.2%, and the lower limit of quantitation was 1% in hydrogenated vegetable oils.

Further refinement of this procedure by means of single-bounce, horizontal attenuated total reflection (SB-HATR) infrared spectroscopy was recently reported.
Using this procedure, only 50 µL (about 2–3 drops) of neat oil (or esters) is placed on the horizontal surface of the zinc selenide element of the SB-HATR infrared cell. The absorbance values obtainable are within the linearity of the instrument. The test portion of the neat oil can easily be cleaned from the infrared crystal by wiping with a lint-free tissue before the next neat sample is applied. The method is accurate for trans concentrations greater than 1%. The SB-HATR FTIR procedure was used to determine the trans content of 18 food products [32]. Recently, this procedure has been collaboratively studied and good statistical results were obtained. For triacylglycerols, the reproducibility relative standard deviations were in the range 18.97–1.62% for 1.95–39.12% trans (as trielaidin) per total fat. For fatty acid methyl esters, the corresponding values varied from 18.46% to 0.9% for 3.41–39.08% trans (as methyl elaidate) per total fat. At worst, accuracy was −11% and averaged 1% low bias.

The ATR-FTIR procedure was voted official method AOCS Cd 14d-99 by the AOCS in 1999 [34] and official method 2000.10 by AOAC International in 2000 [35] after testing in a 12-laboratory international collaborative study. Analytical ATR-FTIR results exhibited high accuracy relative to the gravimetrically determined values. Comparison of test materials with similar levels of trans fatty acids indicated that the precision of the current ATR-FTIR method was superior to those of the two most recently approved transmission infrared official methods: AOAC 965.34 [36] and AOAC 994.14 [37].

The ATR-FTIR method was also evaluated for use with matrices of low trans fat and/or low total fat contents such as milk [38] and human adipose tissue [39]. Preliminary results indicated that the presence of low levels (<1%) of conjugated cis/trans dienes, with absorbance peaks near 985 and 947 cm⁻¹, interfered with the accurate determination of total isolated trans fatty acids [38,39]. Attempts to eliminate interfering absorbance peaks by use of spectral subtraction techniques [38,39] were not satisfactory. In order to overcome the effect of interferences, the ATR-FTIR

Figure 2  IR absorption bands for 0.91–45.87% trans when neat mixtures of methyl elaidate (ME) in methyl oleate (MO) were “ratioed” against neat MO using an ATR liquid cell. (From Ref. 30.)
method was modified by inclusion of the standard addition technique [40]. This modification was applied to several food products, namely, dairy products, infant formula, and salad dressing. This successfully eliminated the interfering absorbance peaks. The presence of <1% CLA in two butter and two cheese products containing 6.8%, 7.5%, 8.5%, and 10.4% trans fatty acids (as a percentage of total fat) would have resulted in errors of −11.6%, 10.4%, 17.6%, and 34.6%, respectively, in trans fat measurements using the unmodified method.

B. Raman Spectroscopy

The use of Raman spectroscopy for the analysis of lipids has been limited by high instrument costs and technical hurdles. However, there have been several developments that should increase its future use, including new detectors and the availability of more powerful computers at reduced cost [41]. Raman spectroscopy seems appropriate to use for the simultaneous analysis of the cis and trans content of hydrogenated oil, since cis monoene, diene, and triene C=C stretch features were found near 1654, 1657, and 1655 cm$^{-1}$, respectively [42]. Also, trans monoene and diene C=C stretch features were found at 1668 and 1670 cm$^{-1}$, respectively [42]. The cis/trans isomer ratios were determined with a precision of 1%. The C=C stretch and the CH$_2$ scissor intensity ratios were also used to determine iodine values. The total unsaturation in neat oils and margarines was determined by FT–Raman spectroscopy [43]. FT–Raman analysis produced significantly improved spectra in one-fifth of the time required for dispersive Raman spectra. Rapid developments in FT–Raman instrumentation indicates that use of this technique will continue to increase.

C. High-Resolution $^{13}$C NMR Spectroscopy

NMR spectroscopy has several applications in the study of lipids. Pulsed low-resolution NMR has been used for the determination of solid fat content, polymorphism, oil content, and moisture in oil, as well as for studying fatty emulsions [44]. High-resolution $^{13}$C NMR spectroscopy also has several applications in fat analysis and is an effective tool that can provide structural information about cis and trans isomers of fatty acid. $^{13}$C NMR spectra exhibit chemical shifts as a series of sharp lines that are measured in parts per million relative to a standard, with the line height being indicative of the intensity [45].

The signal from a carbon that is allylic to a double-bond carbon provides evidence concerning the cis or trans nature of the double bond. The intensity of the signal can also be used for quantitation. The number, position, and geometric configuration of double bonds in the fatty acid chain determine the chemical shifts and intensities of the peaks observed. A summary of the published $^{13}$C NMR chemical shifts [46] for allylic carbons of linoleic acid geometric isomers is shown in Table 1. These chemical shifts are unique and do not overlap with lines for other carbons in the fatty acid chain and, therefore, can be used for identification. For example, in the case of the cis-9,trans-12 isomer, the C-8 carbon allylic to a cis bond has a chemical shift of 27.15 ppm; the C-14 carbon allylic to a trans bond has a chemical shift of 32.6 ppm; and the methylene C-11 carbon allylic to both a cis and a trans bond has a chemical shift of 30.5 ppm. High resolution $^{13}$C NMR was used to determine the total trans-monoene, diene, and triene isomer content of shortening.
Table 1  $^{13}$C NMR Shifts for Allylic Carbons of Linoleic Acid Geometric Isomers

<table>
<thead>
<tr>
<th>Isomer</th>
<th>Chemical shift (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C11</td>
</tr>
<tr>
<td>$^\text{trans}$-9,$^\text{trans}$-12 18:2</td>
<td>35.7</td>
</tr>
<tr>
<td>$^\text{cis}$-9,$^\text{trans}$-12 18:2</td>
<td>30.5</td>
</tr>
<tr>
<td>$^\text{trans}$-9,$^\text{cis}$-12 18:2</td>
<td>30.55</td>
</tr>
<tr>
<td>$^\text{cis}$-9,$^\text{cis}$-12 18:2</td>
<td>25.75</td>
</tr>
</tbody>
</table>

*Source* Ref. 39.

[47]. This technique was also used to confirm the presence of specific trans diene isomers in hydrogenated soybean oil [48].

The $^{13}$C NMR signal from olefinic carbons yields bands that are indicative of the double-bond position in unsaturated oils. Olefinic signals from 128 to 132 ppm provide the most useful information about bond position [44]. Both signals obtained for the two olefinic carbons in cis and trans monoene fatty acids, as well as their differences, can be used to determine double-bond position.

D. FTIR–Partial Least Squares (PLS) Regression

To evaluate nutritional properties of foods, quantitative analysis of trans unsaturation was investigated by using transmission FTIR–partial least squares (PLS) [49]. This approach offered significantly reduced analysis time compared with conventional GC methods and was effective for a wide range of trans values. Improved results were reported for raw materials and food products [49].

A heated SB-HATR sampling accessory was developed and used to develop a method for the simultaneous determination of iodine value (IV) and trans content for neat fats and oils [50]. PLS regression was employed for the development of the calibration models, and a set of nine pure triacylglycerol test samples served as the calibration standards. Satisfactory agreement (SD < 0.35) was obtained between the predictions from the PLS calibration model and trans determinations by the FTIR/SB-HATR method.

E. Near-Infrared Spectroscopy

A generalized partial least squares calibration was developed for determination of the trans content of edible fats and oils by Fourier transform near-infrared (FT-NIR) spectroscopy [51]. The trans reference data, determined by using the SB-HATR mid-IR official method, were used in the development of the generalized FT-NIR calibration. The FT-NIR trans predictions obtained using this generalized calibration were in good agreement with the SB-HATR results. The authors concluded that FT-NIR provided a viable alternative to the SB-HATR [51].

III. ANALYSIS USING CHROMATOGRAPHY

The chromatography of lipids involves the separation of individual components of a lipid mixture as they pass through a medium with a stationary matrix. This matrix
may be packed or bound to a column, as is the case in GC and HPLC, or bound to a glass plate as in the case of thin-layer chromatography (TLC). The mobile phase in gas chromatography is usually an inert gas such as helium or nitrogen. For HPLC or TLC, the mobile phase may be an aqueous or organic solvent. Intact triacylglycerols or fatty acids can be separated, but many chromatography applications consist of separating the methyl ester or other derivatives of individual fatty acids.

A. Gas Chromatography

The most significant occurrence in the chromatography of lipids has been the development of gas chromatography. Fatty acid methyl esters, including saturates and unsaturates, were first successfully separated in 1956 with a 4-foot-long column packed with Apiezon M vacuum grease [52]. Separations were demonstrated based on chain length, degree of unsaturation, and cis or trans geometric configuration.

1. Separations Using Packed Columns

In general, retention times of fatty acid derivatives on nonpolar columns are based on volatility and, therefore, separation occurs primarily by carbon chain length. Retention times of fatty acid derivatives on polar columns are mainly determined on the basis of polarity and chain length. These columns, therefore, are more effective at resolving unsaturated fatty acids with different degrees of unsaturation. The chromatographic characteristics of several packed columns (Silar 10C, Silar 9CP, SP 2340, and OV-275), made available in the mid-1970s as a result of the then recently developed silicone stationary phases, were compared [53]. The development of these highly polar, temperature-resistant columns made it possible to separate the geometric isomers of fatty acids. A column packed with 12% Silar 10C as the stationary phase was able to baseline separate the all-cis from the all-trans components of octadecadiynelic fatty acid methyl esters. All eight of the geometric isomers of linolenic acid were also partially resolved. However, it was not possible to effectively separate individual positional cis and trans isomers using a 20-foot-long packed column [48]. This drawback of packed columns was due largely to practical considerations that limited separation efficiencies, even with highly selective stationary phases [54]. Although larger numbers of theoretical plates could be produced, the need to compensate for back pressure prevented the operation of these columns at more than 6000 theoretical plates. However, a packed column was successfully used to determine trans concentration in a collaborative study [55]. This study concluded that GC analysis using an OV-275 packed column was as effective as IR spectroscopy in quantitating total trans unsaturation.

2. Separations Using Capillary Columns

The development of flexible fused silica columns in the early 1980s [56] led to the popularity of capillary columns, which dramatically increase the number of effective plates, thus improving column separation efficiencies. The number of effective plates could be raised from 40,000 to 250,000 by increasing the column length from 15 m to 100 m.

The increased availability of long capillary columns with a variety of diameters and stationary phases has decreased the use of packed columns in the last 10 years. As Figure 3 indicates, several monoene, diene, triene, and conjugated fatty acid
methyl ester isomers from partially hydrogenated soybean oil were separated in a single run on a 100-m capillary column [57]. Several individual trans isomers appeared to be baseline-resolved by capillary GC. When GC conditions were optimized to separate trans diene methyl esters, near-baseline resolution was obtained for several of these isomers, as shown in Figure 4 [58]. The trans triene isomers present in partially hydrogenated soybean oil were separated into four peaks on a 100-m capillary GC column [59]. Fractional crystallization and reversed phase HPLC, followed by GC analysis of hydrazine reduction products, served to identify and quantify these triene isomers.

Most laboratories use capillary columns to quantitate individual fatty acid isomers found in hydrogenated oils. The application of capillary GC analysis in the separation and identification of positional and geometric isomers of unsaturated fatty acids has been reviewed [60]. This analysis becomes even more complex for geometrical and positional isomers found in hydrogenated fish oils, which contain many more fatty acid isomers than are present in hydrogenated vegetable oils. An extensive analytical study of hydrogenated menhaden oil using capillary GC was conducted [61,62]. At an iodine value of 84.5, the most unsaturated isomers were eliminated, although 13.1% diene, 8.3% triene, and 0.4% tetraene isomers were still present. The 20-carbon isomers were analyzed further to determine the cis and trans bond positions. A wide range of monoene, diene, and triene cis and tran positional isomers were identified at virtually every position of the fatty acid chain.

The content of trans-C16:1 in human milk, as determined by GC, is usually high due to the overlap with peaks attributed to C17 fatty acids [63]. Isolation by
Figure 4  The C-18 region of the gas chromatogram of the fatty acid methyl esters of canola–palm–butter spread using an SP-2560 100 m × 0.25 mm fused silica capillary column. Peak identification: 12,14,16–18, t.t-NMID; 19, 9,t12; 20, c9,t13; and t8,c12; 21, t.tc; 22, t8,c13?, and cc-MID; 23, c9,t12; 24, c8,c13?; 25, 9,c12; 26, 9,c15, and t10,c15?; 27, c9,c13?; 28, c9,c12; 29, c9,c15. NMID, non-methylene-interrupted diene; MID, methylene-interrupted diene. (From Ref. 58.)
in the transition periods in spring and late autumn. The average contents were \( \text{c9t11-18:2} \) 0.76%, \( \text{(trans-C18:1)} \) 3.67%, \( \text{(trans-C18:2)} \) 1.12%, and (total trans fatty acids) 4.92%. High correlation coefficients \((r)\) were reported between the content of the CLA isomer c9t11-18:2 and the contents of trans-C18:1, trans-C18:2, total trans fatty acids, and C18:3 of 0.97, 0.91, 0.97, and 0.89, respectively. A probable metabolic pathway in the biohydrogenation of linolenic acid was also reported: \( \text{c9c12c15} \rightarrow \text{c9t11c15} \rightarrow \text{t11c15} \rightarrow \text{t11} \) [65].

3. Equivalent Chain Length

It is difficult to identify peaks in gas chromatograms, mainly because of the complexity of the fatty acid composition of hydrogenated vegetable and fish oils. Some of the problems of using capillary GC for qualitative and quantitative analysis of hydrogenated oils have been reviewed [56]. The major identification problem is a result of the large number of fatty acid isomers present. One difficulty in identifying peaks is due to the fact that some polyunsaturated fatty acids can have longer retention times than the next longer chain fatty acid. This happens most frequently on highly polar columns used to separate fatty acid isomers.

One way to predict where a certain fatty acid will elute on a GC column is based on the so-called equivalent chain length (ECL). The use of ECL to identify fatty acid isomers on GC columns has been reviewed [56,66]. It was reported that ECLs are constants for a specific carrier gas and column and are independent of experimental conditions [67]. A specific fatty acid can be characterized by obtaining its ECL on both polar and nonpolar columns. The ECL consists of one or two integers indicating the positional chain length, and two numbers after the decimal that indicate the fractional chain length (FCL). The ECL values for fatty acids on a specific column are determined by first using semilog paper to plot the log of the retention times (y axis) against those of the saturated fatty acids (e.g., 16:0 = 16.00, 18:0 = 18.00, 20:0 = 20.00) on the x axis observed under isothermal conditions. Using the same column, the retention time (y-axis value) of an unsaturated fatty acid is then plotted to find its ECL on the x axis. For example, the ECL of linoleic acid \( (18:2n-6) \) was found to be 18.65. The FCL value was 0.65 (18.65 − 18.00). Using these data, the ECL for 20:2n-6 was predicted to be 20.65 (20.00 + 0.65). The actual ECL of 20:2n-6 was experimentally determined to be 20.64 [36]. Equivalent chain length data from a GC analysis of rapeseed oil showed that the cis-11 20:1 fatty acid coelutes with trans triene isomers at oven temperatures above 160°C [68].

4. Relative Response Factors

Area normalization is often used to report the relative concentration of fatty acid isomers in an oil mixture. This method is accurate only if all isomers exhibit the same response in the GC detector. However, all the fatty acid isomers do not give the same relative response in the GC flame ionization detector. Relative response factors, first proposed in 1964 [69], were found to depend on the weight percent of all carbons except the carbonyl carbon in the fatty acid chain. A later study confirmed these findings and concluded that the required corrections were not insignificant, even though they were often not used [70]. Failure to use a correction factor could result in an error of about 6% (relative to 18:0) in the case of 22:6. Table 2 summarizes some of the correction factors recommended.
Table 2  Response Factors of Unsaturated Esters Relative to Methyl Stearate

<table>
<thead>
<tr>
<th>Fatty acid methyl ester</th>
<th>Response factor</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>18:0</td>
<td>1.000</td>
<td>—</td>
</tr>
<tr>
<td>18:1</td>
<td>0.996</td>
<td>0.003</td>
</tr>
<tr>
<td>18:2</td>
<td>0.986</td>
<td>0.001</td>
</tr>
<tr>
<td>18:3</td>
<td>0.981</td>
<td>0.003</td>
</tr>
<tr>
<td>20:4</td>
<td>0.959</td>
<td>0.005</td>
</tr>
<tr>
<td>22:6</td>
<td>0.941</td>
<td>0.002</td>
</tr>
</tbody>
</table>

Source  Ref. 70.

B. Thin-Layer Chromatography

Thin-layer chromatography (TLC) has been widely used to separate classes of lipids on layers of silica gel applied to glass plates. TLC is simple to use and does not necessarily require sophisticated instrumentation. In most cases, TLC using silica gel does not separate fractions on the basis of number or configuration of double bonds in fatty acid mixtures [71]. The most effective TLC separation of cis and trans isomers has been achieved by means of argentation: layers of silica impregnated with silver nitrate.

Argentation is the general term used to describe methods that use the long-known principle that silver ion forms complexes with cis more strongly than with trans double bonds. In argentation chromatography the separation of cis and trans isomers depends on the relative interaction strength of the $\pi$ electrons of double bonds in each isomer with silver ions. The actual interaction of each fatty acid isomer depends on the geometry, number, and position of double bonds. Argentation TLC (Ag-TLC) has been used for several years to separate unsaturated lipids on thin-layer plates [72].

The use of argentation methods, including TLC, countercurrent distribution (CCD), and liquid column chromatography, has been reviewed [72,73]. Several classes of fatty acid isomers can be isolated using preparative argentation TLC. These isolated fractions can then be analyzed further by using ozonolysis [61,62,74–76] and GC. Good resolution can be achieved for a variety of fatty acid methyl esters using two-dimensional argentation TLC separation and reversed phase chromatography on a single plate [77,78].

Incorporating silver ions with the silica gel on TLC plates [78] creates a number of disadvantages: (1) large amounts of expensive silver nitrate are required; (2) the method is labor-intensive; (3) the separated components can be difficult to visualize; (4) autoxidation can occur on the plate; and (5) some silver ions may elute with the fractions. Silver ion TLC was used to completely fractionate the trans and cis monoenes [80,81]. The Ag-TLC fractions were then analyzed by capillary GC. The total trans content of a partially hydrogenated vegetable oil was obtained by argentation TLC followed by GC analysis [29,58].
C. High-Performance Liquid Chromatography

For most applications, HPLC does not exhibit the separation efficiencies offered by long capillary GC columns. It does, however, offer several advantages. For example, the mild conditions used in HPLC work enable heat-sensitive components to be separated, a feature that reduces the possibility of isomerization taking place during the analysis of unsaturated fatty acids. Another advantage is that fatty acid fractions can be collected and analyzed further using hyphenated techniques such as GC-FTIR and GC-MS.

A variety of detectors are used to identify lipid solutes as they elute from HPLC columns. The most common HPLC detector has been ultraviolet-based equipment that allows one to monitor the column effluent at about 200 nm. The lack of strong absorbing lipid chromophores limits the sensitivity of this detection method unless fatty acid derivatives that absorb strongly in the UV range are prepared. Refractive index (RI) detectors are also commonly used but cannot be paired with solvent gradients and are sensitive to temperature fluctuations. Evaporative light-scattering detectors (ELSD) can be used with solvent gradients, are at least as sensitive as the best RI detectors, and can be used for quantitation. The advantages of these and other detectors useful for HPLC analysis have been reviewed [82].

1. Reversed Phase HPLC

Fatty acids and fatty acid methyl esters were successfully separated on Vydac reversed phase columns in an early study [83]. In another study, a Bondapack C18 reversed phase column was used with a methanol–water mobile phase to separate methyl elaidate and oleate [84]. Five fractions from hydrogenated menhaden oil, containing saturated fatty acids as well as cis and trans monoene isomers, were collected with two C18 preparative HPLC reversed phase columns (Waters, 5.7 cm × 30 cm) connected in series [85].

2. Silver Ion HPLC

The many advantages of using argentation HPLC columns over TLC plates [84] include reproducible separation of analytes, column reusability, short run times, and high recoveries. Until recently, a lack of stable columns with controlled silver levels limited the use of silver ion HPLC separations [71]. The silver ions would bleed from the silica adsorbent, cause unpredictable results, shorten the column life, and hinder the reproducibility of the results. A more successful argentation HPLC procedure was developed by linking the silver ions via ionic bonds to a silica–phenylsulfonic acid matrix [86]. This column gave excellent reproducible separations for triacylglycerols, fatty acids, and their positional and geometric isomers [87]. This column was also used for the separation, collection, and quantification of all eight geometric isomers of linolenic acid phenacyl esters, with a mobile phase ranging from 5% methanol in dichloromethane to a 50:50 solvent mixture [88]. Figure 5 shows the excellent resolution obtained for 18:3 isomers using the silver ion HPLC column.

Commercial silver ion HPLC columns were recently introduced and should dramatically increase the use of this technique. These columns have been reported to give satisfactory results for up to one year [71]. A commercially available (Chrom-pack Ltd.) argentation HPLC column with an acetonitrile–hexane mobile phase was
used to separate the cis/trans fatty acid isomers of methyl oleate, methyl linoleate, methyl linolenate, and 15 of the 16 cis/trans methyl arachidonate positional isomers [89]. In other research, this column was used with a 0.15% acetonitrile in hexane isocratic mobile phase to obtain four fractions from hydrogenated vegetable oil that were subsequently analyzed by capillary GC [90]. As Figure 6 shows, the silver ion HPLC column separated the cis and trans monoene positional isomers of fatty acid methyl esters with 0.08% acetonitrile in hexane mobile phase. Quantitation of positional isomers was obtained with this silver ion HPLC procedure. Comparable quantitation was also achieved with a 100-m capillary GC column used to analyze the isolated trans HPLC fraction.

IV. ANALYSIS USING COMBINED TECHNIQUES

The direct quantitation of all fatty acid isomers in hydrogenated oils by gas chromatography is nearly impossible. There is overlap of cis/trans monoene and diene positional isomers even with the high efficiency of long capillary columns [68,89–91]. Several analytical techniques have been combined with gas chromatography to determine accurate fatty acid profiles of hydrogenated oils including ozonolysis and infrared spectroscopy.

A. Ozonolysis/Gas Chromatography

Ozonolysis followed by gas chromatography can be used to determine the double-bond position of cis and trans isomers. Ozonolysis cleaves the hydrocarbon chain of fatty acids at positions of unsaturation. This includes reductive cleavage to form aldehydes, aldehyde esters, and other fragments, which are identified according to their retention times on GC columns compared with known standards. Ozonolysis
Figure 6  Silver ion HPLC chromatogram of partially hydrogenated vegetable oil methyl ester 18:1 positional isomers. Mobile phase: 0.08% acetonitrile in hexane. Fractions: A, saturates; B, trans-18:1; C, cis-18:1. (From Ref. 90.)

has been used to determine the double-bond position of both monoene [74,92] and diene isomers [93,94]. Even though the double-bond position can be determined by ozonolysis and GC, the geometric configuration cannot. Computer programs involving solutions of several simultaneous equations were used to calculate fatty acid bond position from ozonolysis results [94]. A recent review discusses the use of ozonolysis and other chemical methods for lipid analysis [95].
B. Gas Chromatography/IR Spectroscopy

GC provides useful information concerning the total fatty acid composition of hydrogenated oils. The overlap of some of the \textit{trans} with \textit{cis} monoene peaks makes it difficult to get accurate GC determinations of total \textit{cis} and total \textit{trans} monoene content of food products. Accurate determination of these isomers is essential because the latest food labeling regulations (Nutrition Labeling and Education Act (NLEA) [96]) permit \textit{cis} monounsaturates to be listed on a food label. A combined GC and IR method to determine \textit{cis} and \textit{trans} fatty acid monoenes that coelute on GC columns was developed [68]. This method was adopted as an AOAC official method [97] upon completion of a collaborative study [57]. In this method, the total \textit{trans} content of the oil was first determined by modifying a published IR procedure [23] that used a two-component calibration plot. Then the weight percentages of all the \textit{trans} diene (18:2\textit{t} and 18:2\textit{tt}) and triene (18:3\textit{t}) isomers were determined by means of a highly polar 50- or 100-m capillary GC column. The weight percentage of \textit{trans} monoenes (18:1\textit{t}) was then determined by the following formula with the appropriate correction factors (0.84 and 1.74):

\[
\text{IR } \text{trans} = \frac{\%18:1t}{100} + (0.84 \times \%18:2t) + (1.74 \times \%18:2tt) + (0.84 \times \%18:3t) \tag{5}
\]

After the \textit{trans} monoenes had been calculated, the \textit{cis} monoenes were determined by finding the difference between the total monoenes determined by GC and the \textit{trans} monoenes that were calculated:

\[
\%18:1c = (\text{total } \%18:1t \text{ by GC} + 18:1c \text{ by GC}) - (\%18:1t \text{ by calculation}) \tag{6}
\]

C. Silver Ion Chromatography/NMR

As already discussed, $^{13}$C NMR spectroscopy can be an effective tool for the identification of fatty acid isomers. This tool is most useful if purified fatty acid fractions are first obtained. Silver ion chromatography on both TLC plates and HPLC columns have been used to obtain purified fractions for $^{13}$C NMR analysis. Hydrogenated soybean oil was separated into six bands with preparative silver ion TLC [48]. The isomers in each band were identified by capillary GC and $^{13}$C NMR analysis. The presence of specific \textit{trans} diene isomers was confirmed by observing their unique chemical shifts. Recently, silver ion HPLC was used to obtain a purified \textit{trans} monoenone fraction from hydrogenated soybean oil [98]. $^{13}$C NMR spectroscopic analysis of this fraction confirmed the presence of the minor $\Delta 6$ and $\Delta 7$ \textit{trans} monoene positional isomers. The presence of the $\Delta 5$ \textit{trans} monoene positional isomer was inferred. This HPLC fraction had been analyzed by capillary GC (see Sec. V.B), but several minor isomers could not be identified.

V. ANALYSIS USING HYPHENATED TECHNIQUES

The lack of standards for many fatty acids and their isomers and the problem of coelution can lead to the misidentification of fatty acids in gas chromatograms. Recent publications misidentified \textit{trans} monoene positional isomers [90,98]. Another study determined the level of \textit{trans},\textit{trans} 18:2 isomers in margarine to be an order of magnitude too high (3\% instead of 0.3\%) also because of GC peak misidentification.
cation [100]. Similarly, the published literature contains an incorrect report that liquid canola shortening was contaminated with fatty acids found in animal fat [101].

To help identify peaks, a GC column can be interfaced to another instrument such as an infrared spectrometer or a mass spectrometer. Hyphenated techniques use on-line detection to confirm the identity of peaks in a chromatogram. Griffiths et al. [102] have reviewed interfaces between gas, supercritical fluid, and high performance liquid chromatography and Fourier transform spectrometry (GC-FTIR, SFC-FTIR, and HPLC-FTIR, respectively). Some of these hyphenated techniques were first used to elucidate the structure of unusual fatty acid isomers, including cyclic fatty acid monomers (CFAM) that often contain trans double bonds in the hydrocarbon chain and cis double bonds in five- or six-membered rings. The formation and the biological effects of these cyclic compounds have been reviewed [103].

A. GC-FTIR

For many GC-FTIR instruments, the effluent from the GC column flows continuously through a light pipe (LP) gas cell [102]. LP instruments generally have detection limits of 10–50 ng for unknown complex mixtures of fatty acid methyl esters. On-line infrared spectra of CFAM peaks eluting from a gas chromatogram were obtained with an LP GC-FTIR [104]. The results were used to show which of these CFAM contained cis and/or trans double bonds. Several minor peaks in this mixture could not be identified because of the limited sensitivity of the method.

Bourne et al. [105] developed an improved GC-FTIR technique using a matrix isolation (MI) interface that increases the sensitivity of the GC-FTIR determination by an order of magnitude. GC–matrix isolation–FTIR is extremely useful for quantitating peak area, determining peak homogeneity, and obtaining structural information on a compound. Although the LP GC-FTIR technique can confirm the identity of intact molecules, its detection limits are higher than those of other GC detectors (e.g., flame ionization). Matrix isolation (MI) is a technique in which analytes and an inert gas (argon) are rapidly frozen at cryogenic temperatures (12 K) and are trapped as a solid matrix on the outer rim of a moving gold-plated disk. The IR spectra of these molecules are free from indications of intermolecular hydrogen bonding and other band broadening effects. These combined benefits yield greater sensitivity that equals, for many applications, that of gas chromatography–mass spectrometry (GC-MS) [106].

GC-MI-FTIR was used to quantitate low levels of saturated trans monoene [85], diene [107,108], triene, and conjugated diene fatty acids and their isomers [108] in hydrogenated vegetable and/or fish oils. Figure 7 shows the IR spectra (out-of-plane deformation absorption bands) for two trans-conjugated diene geometric isomers. The characteristic absorption bands shown were used for identification. The conjugated trans,trans diene isomer had an out-of-plane deformation absorption band at 990 cm⁻¹, whereas, the cis,trans isomer had absorption bands at 950 and 986 cm⁻¹. The absorption band for the corresponding methylene-interrupted trans diene and the trans monoene was at 971 cm⁻¹. There were also unique absorption bands for trans and cis diene isomers in the area of the spectrum typical of the carbon–hydrogen stretch vibrations (3035/3005 cm⁻¹ and 3018 cm⁻¹, respectively) and in the area typical of the carbon–hydrogen out-of-plane deformation absorption bands (972 and 730 cm⁻¹, respectively).
Figure 7  Expanded IR spectral range showing out-of-plane deformation bands for conjugated cis-trans (top spectrum) and trans-trans (bottom spectrum) 18:2 dienes. (From Ref. 108.)

GC-MI-FTIR is also effective in determining the concentration of individual fatty acid methyl esters without having to consider the relative response of the gas chromatograph’s flame ionization detector (FID). It is possible to quantitate trans isomers even with partial GC peak overlap from cis isomers. Quantitation of trans diene isomers was based on measurement of the height of the observed C—H out-of-plane deformation band at 971 cm$^{-1}$ for trans groups and that of the CH$_3$ asymmetric stretching band at 2935 cm$^{-1}$ for the 17:0 internal standard [107]. Calibration plots of absorbance versus nanograms injected were generated for the range of 2–33 ng. Recovery (on cryogenic disk) was based on the determination of the internal standard. The amount of analyte present in injected aliquots was calculated from the observed absorbance values and the corresponding calibration plot.
When GC-MI-FTIR was used to quantitate the trans monoene isomers in margarine, the results had a high bias relative to the GC FID’s response [107,108]. The higher GC-MI-FTIR values presumably resulted from its higher specificity. This is because the MI-FTIR determination is based on a discriminatory feature (971 cm⁻¹ absorbance band) that is observed only for trans species. The intensity of this band is not affected by cis isomers even when they chromatographically overlap. These results confirm the conclusion that quantitation by means of GC peak areas can result in an underestimation of trans monoenes.

GC-FTIR analysis was used to determine the geometric configuration of fatty acid methyl esters separated by silver ion HPLC [110]. This was achieved with the more recent direct deposition (DD) interface that condenses GC eluates on a moving zinc selenide window cooled to near liquid nitrogen temperature. This DD instrumentation is even more sensitive than the matrix isolation interface [111] because the analytes are condensed on a track that is about 100 μm wide, and microscope objectives are used to collect and focus the infrared beam. Unlike GC-MI-FTIR, during GC-DD-FTIR operations, the analytes are not diluted in an argon or any other matrix.

B. GC-EIMS

MS can be a very effective tool when used in combination with GC to determine the location of double bonds in fatty acids and their positional isomers. The major problem of analyzing mass spectra of fatty acid methyl esters is the tendency of the double bonds to migrate during electron ionization. The mass spectra exhibit low mass ions that do not provide structural information. Some of the methods used to overcome this problem include soft ionization and derivatization [112]. Chemical ionization (CI) methods for the determination of double- and triple-bond positions were reviewed [113]. A CI-MS procedure was used to determine the double-bond positions in fatty acids from marine organisms [114].

The most successful approach has been to derivatize the carboxyl group to a nitrogen-containing compound. Common derivatizing agents have included pyrrolidine, picolinyl ester, and 4,4-dimethyloxazoline (DMOX). A recent review of these derivatives indicates that the most useful ones by far are the picolinyl ester and DMOX derivatives [115]. With these derivatives, double-bond ionization and migration are minimized. Simple radical-induced cleavage occurs at each C—C bond along the chain. Therefore, for unsaturated fatty acids containing up to several double bonds, there is decreased abundance of low mass ions and an increase in a series of ions resulting from carbon—carbon bond scission. Diagnostic ions occur wherever there is a functional group in the chain that interrupts the pattern of cleavage from C—C bonds. A C==C bond or a five- or six-membered ring might be responsible for such disruptions.

GC-EIMS analysis of picolinyl ester derivatives can unequivocally identify polyunsaturated fatty acids [116,117]. In the work just cited, however, the GC resolution of the picolinyl fatty acid esters was not as good as that of other derivatives. Reversed phase HPLC fractionation of the picolinyl fatty acid esters prior to identification by GC-MS was necessary to obtain acceptable results for hydrogenated samples [118]. A total of 39 fatty acid components in cod liver oil were identified using this method. In a more recent study, silver ion HPLC was used to fractionate
CFAMs before converting them to the picolinyl ester derivatives [119]. Some of the GC problems associated with picolinyl esters could be overcome by using high temperature, low bleed, cross-linked polar columns [120].

Mossoba et al. reported that 2-alkenyl-4,4-dimethyloxazoline derivatives of diunsaturated CFAMs exhibited distinctive mass spectral fragmentation patterns that could be used to pinpoint the positions of double bonds and of 1,2-disubstituted, unsaturated, five- and six-membered rings along the hydrocarbon chain [121]. One of the advantages of using the DMOX derivatives was the good chromatographic resolution: sometimes higher than that observed for derivatives of fatty acid methyl esters [122,123]. Most CFAMs in heated flaxseed oil were identified, and the double-bond configurations (cis or trans) were unequivocally established by using GC-MI-FTIR (Table 3).

Recently, Mossoba et al. [110] confirmed the identity of individual trans monounsaturated fatty acid positional isomers in partially hydrogenated soybean oil. Fatty acid methyl esters were fractionated by silver ion HPLC and then analyzed by GC-DD-FTIR to determine geometric configuration and by GC-EIMS on DMOX derivatives to determine double-bond position [110]. GC peak resolution obtained with a 100-m capillary column was higher for the DMOX derivatives than for those of the fatty acid methyl esters. Figure 8 demonstrates the excellent resolution that was obtained for DMOX derivatives of trans monoene positional isomers. The bottom GC profile, with about twice the amount of injected sample, shows evidence of overload in the early part of the trace but enhanced response for the $\Delta 13$ and $\Delta 14$ positional isomers. The double-bond positions for nine individual trans monoene positional isomers were confirmed by their unique DMOX mass spectra. Most significantly, this was the first report of the capillary GC separation of the $\Delta 13$ and $\Delta 14$ trans monoene positional isomers in hydrogenated vegetable oil. Figure 9 presents the mass spectral data that confirmed their bond position. The identity of the trans-$\Delta 13$ isomer was further confirmed by comparison with a standard. Hence, the double-bond position and configuration could be readily established for a complex distribution of monounsaturated fatty acid isomers found in dietary fat by using the two hyphenated techniques, GC-DD-FTIR and GC-EIMS.

C. SFC-FTIR

Capillary SFC is an effective tool for separating nonpolar to moderately polar complex mixtures of natural products having molecular weights of 100–1000 Da. SFC uses a gas compressed above its critical temperature and pressure to carry analytes through a chromatographic column. The many applications of SFC to the analysis of lipids have been reviewed [124–128]. SFC can be used to separate some compounds at lower temperatures than GC, and a simultaneous interface with FID and FTIR detectors can be readily achieved. Different SFC columns can be used to separate lipids according to carbon number or degree of unsaturation. However, capillary GC and silver ion HPLC columns give better peak resolution when separating fatty acid derivatives from hydrogenated oils.

SFC has been used to obtain structural information, backed up by mass spectrometry for detection. The main limitation of early applications has been the lack of suitable commercial interfaces [129].
<table>
<thead>
<tr>
<th>GC peak</th>
<th>Ring, five-membered</th>
<th>Chain, trans</th>
<th>Ring, six-membered</th>
<th>Chain, cis</th>
<th>Chain, trans</th>
<th>Chain, trans</th>
<th>Ring, Six-membered</th>
<th>Ring, Five-membered</th>
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*Source* Ref. 123.
Microscale supercritical fluid extraction (SFE) can be directly coupled to a capillary SFC column. This procedure was used to analyze the fatty acid composition of a 1-mg sample of cottonseed kernel [128].

SFC with an FTIR detector (SFC-FTIR) has been used to elucidate structural information from fatty acid isomer mixtures. Standards as well as fatty acids were separated from coconut oil, Ivory soap, soybean oil, and butterfat by means of a packed SFC column [130]. The presence of unsaturated fatty acids was verified by an IR absorption maximum at 3016 cm\(^{-1}\). SFC-IR spectroscopy was used to analyze triacylglycerols and free fatty acids under similar conditions [131]. One study featured an SFC coupled to an FTIR spectrophotometer equipped with a LP flow cell to determine the level of trans unsaturation in partially hydrogenated soybean oil [132]. The IR absorption band at 3016 cm\(^{-1}\) indicated the presence of linoleic acid, while the 972 cm\(^{-1}\) absorption band indicated the presence of trans unsaturation.

A packed microcolumn argentation SFC system was used to separate and quantify triacylglycerols in vegetable, fish, and hydrogenated oils [133]. This system gave excellent separations for these complex samples and was as effective as HPLC.
**Figure 9** GC-EIMS spectra for the Δ13 and Δ14 pair of *trans* 18:1 DMOX positional isomers; the ions due to allyllic cleavage are at the following positions: Δ13, 238, 293, and 306 nm; Δ14, 252, 306, and 320 nm. (From Ref. 110.)

### D. LC-MS

HPLC is the method of choice for the determination of both neutral and polar glycerol lipids. HPLC coupled with mass spectrometry (LC-MS) provides separation as well as unequivocal detection of all lipid species [134]. Application of thermospray HPLC was reported for the analysis of long chain carboxylic acids and their methyl esters [134,135]. This technique was paired with reversed phase HPLC to identify standard triacylglycerols and glycerophospholipids.

The use of the electrospray technique with mass spectrometry was recently reviewed [136]. A dramatic increase in the use of this technique to analyze a variety of lipids, including complex mixtures of triacylglycerols, is expected in the near future.

Using reversed phase HPLC coupled with flame ionization detection or quadrupole mass spectrometry with an atmospheric pressure chemical ionization source, the triacylglycerol compositions of margarine base stock products were determined [137]. These were highly-saturated vegetable oils, randomized vegetable oils, vegetable oil–hard stock blends, and interesterified vegetable oil–hard stock blends. Triacylglycerol percent composition results for randomized and interesterified test samples of known composition exhibited less average error using the HPLC quadrupole method, after application of response factors, than the results by HPLC coupled with...
flame ionization detection. The fatty acid compositions calculated from the MS data exhibited less average error than the fatty acid determinations using flame ionization detection data [137].

E. MS-MS

The use of tandem mass spectrometry (MS-MS) has grown considerably in the last 10 years [138,139]. The combination of LC-MS-MS with electrospray ionization is the most effective current technique for lipid ester analysis [134]. While mass spectrometry studies the fragmentation pattern of an ionized molecule, MS-MS studies the fragmentation of a particular fragment ion present in the mass spectrum. Complex mixtures of fatty acids from bacteria were analyzed by means of GC-MS-MS to determine double-bond position [140,141].

In one of the more elegant demonstrations of the potential of combining analytical procedures, silver ion HPLC was used to fractionate the triacylglycerols from winter butterfat. The fractions were subjected to reversed phase HPLC, MS, and MS-MS analysis [142,143]. The distribution of cis and trans isomers within each acyl carbon number of both disaturated monoenoic and saturated dimonoenoic triacylglycerols was elucidated. In future work this procedure could also be used to determine the distribution of trans fatty acids in hydrogenated oils.

VI. trans ISOMERS IN COMMERCIAL PRODUCTS

Although the relationship of dietary lipids to human health is a complicated issue that has sparked some controversy, the importance of controlling fat intake to help maintain an active and healthy lifestyle has been recognized for many years. Consumer health concerns about the types and content of dietary fat has resulted in a great deal of research. trans Fatty acids have received negative publicity in the media with respect to their effects on serum cholesterol. Several food products have been reformulated to increase their unsaturated fat and reduce their total fat and trans fatty acid content.

A. Dietary Recommendations

During the first half of the century, there was little concern about the nutritional properties of dietary fat. This started to change in the 1950s with epidemiological studies, which at the time were controversial, linking diets high in animal fats with the incidence of coronary heart disease [144]. Several studies then indicated the cholesterol-raising effects of saturated fats and the cholesterol-lowering effects of polyunsaturated fats [145,146]. Monounsaturated cis and trans fats were considered neutral. More recently, there have been conflicting reports on the health effects of trans fatty acids, but little cause for concern [4]. An excellent discussion of the dietary recommendations for lipids from several sources, including the National Academy of Science (NAS) and the U.S. Department of Agriculture (USDA), was recently published [147]. Included was a discussion of the Nutrition Labeling and Education Act [96]. The trans fatty acid issue is not addressed by the NLEA, but the regulations authorize the secretary of health and human services to require information on other nutrients to be added to food labels at a later time. Also, the preamble to the regulations indicates that data to determine whether consumers
needed trans fatty acid information to plan healthy diets were insufficient at the time of the act’s passage. The FDA has since been inundated by petitions, nutrition studies, dietary intake studies, and food composition data from groups and individuals on both sides of the trans fatty acid issue.

B. Dietary Intake

There has been little agreement concerning the ideal level of trans fatty acid consumption. One estimate indicated that daily trans consumption could be as high as 38.6 g in some individuals, with an average consumption of about 13 g/day [148]. Subsequent reports, however, suggest that these estimates may have been based on “questionable composition and consumption data,” with the result that the average consumption of trans fatty acids in the U.S. diet may instead be closer to 8 g per person per day [149,150]. The International Life Sciences Institute report [4] cited at the beginning of the chapter summarized consumption data from several studies and found the average daily consumption to range from 2.7 to 12.8 g of trans fatty acids per day. The four studies that differentiated between animal/dairy and vegetable oil sources of trans reported that 78–92% of the trans fat came from vegetable oil sources.

C. trans Content of Food Products

An indicated by the intake data, the main source of fatty acid isomers, including trans fatty acids in the U.S. diet, is the partially hydrogenated vegetable oil used to make margarine and shortening. The positional isomer distribution of the double bonds of trans-octadecenoic acids reported for Canadian margarines ranged from the n-4 to the n-14 position, with most of the isomers at the n-8, n-9, and n-10 positions [151]. It is interesting that different brands of margarine made from the same oil raw materials had significantly different double-bond patterns.

An analysis of over 90 U.S. commercial margarine products in 1985 found that the total amount of trans isomers varied between about 10 and 30% [152]. The trans positional isomers were fairly evenly distributed throughout the fatty acid chain, whereas the cis positional isomers were concentrated at the n-9 and n-10 positions. The diene content of the margarines varied greatly, and all dienes were reported to be 9,12-18:2 isomers [152]. A survey of 197 food products in Germany [153] found the following levels of trans fatty acids:

- Milk and milk products, 1.9–7.9%
- Meat from ruminants, 2–10.6%
- Pork fat, less than 0.5%
- Foods that may contain hydrogenated oils, 0–34.9%

A 1996 capillary GC survey of 18 food products in the United States found that the level of trans relative to the total fat content varied from 1.5% for “fat-free” margarine to 42.9% for chicken-flavored crackers [32].

If, as reported, the level of trans fatty acids in food products has indeed decreased in the last several years, this result is most likely due to improved processing as well as the adverse publicity trans fatty acids have received. For example, a French study reported a decrease of trans monoenes in tub margarines from 13% of the total fat in 1991 to 3.8% in 1995 [154]. However, the study also reported trans
levels in excess of 50% of the total fat for two of three shortenings analyzed. A 1992–1995 Danish study of margarine and shortening [155] also found a decreased level of trans fatty acids. The most significant decrease was from 9.8% trans of the total fat in 1992 compared with 1.2% in 1995 for margarines with a linoleic acid content of 20–40%. There was a 40–90% decrease in trans levels when the 1995 results were compared to an earlier (1985) Danish survey [156].

VII. CONCLUSIONS

As long as the nutritional effects of trans fatty acids are debated, there will be an important need to accurately determine the total trans content of food products by simple and rapid methods. As reviewed in this chapter, the standard infrared methods have been recently revised to increase their accuracy and reliability in the measurement of low levels of trans in food products. Other studies are now in progress to directly determine the trans content of neat fats and oils by means of an infrared single-beam ratioing procedure, with attenuated total reflection cells to increase accuracy and analysis speed.

Also, the nutritional effects of most positional and/or geometric isomers of fatty acid have not been extensively studied. The identification and quantitation of individual fatty acid isomers will be a critical step in any future study designed to determine the nutritional properties of various cis and trans geometric and positional isomers with one or more double bonds. The complex isomer mixture inevitably contains overlapping cis and trans GC peaks that make identification difficult. GC on long, highly polar columns will continue to be an important method for separating fatty acid isomers. However, as discussed in this chapter, GC alone cannot separate all fatty acid isomers in hydrogenated oils. Therefore, combined and/or hyphenated techniques are necessary to separate and identify individual fatty acid isomers. The recent introduction of commercial silver ion HPLC columns should increase the use of these columns, which can be combined with other techniques, such as reversed phase HPLC, SFC, GC, NMR, FTIR, MS, and MS-MS. An increased use of MS-MS in lipid analysis is expected. The introduction of the more sensitive GC-DD-FTIR system and the use of DMOX derivatives, which enhance GC resolution while stabilizing the charge during electron ionization of fatty acids, are other developments that should allow accurate confirmation of individual fatty acid geometric and positional isomers.

Analyzing the total trans content and the individual isomers in complex mixtures in both natural and hydrogenated oils is challenging. With the introduction of some of the new methods discussed in this chapter, there is now an opportunity to meet this challenge. Work on improving these methods is continuing. Although an in-depth discussion of each method is beyond the scope of this chapter, we have attempted to indicate some of the difficulties of analyzing trans fatty acid isomers while referring the reader to some exciting new developments that are now available to lipid scientists. The references provided will help the serious analytical chemist find an in-depth discussion of these methods.

REFERENCES


Chemistry of Frying Oils

KATHLEEN WARNER
U.S. Department of Agriculture, Peoria, Illinois

I. INTRODUCTION
Deep-fat frying is a common method of food preparation that imparts desired sensory characteristics of fried food flavor, golden brown color, and crisp texture. During frying, at approximately 190°C, as oils thermally and oxidatively decompose, volatile and nonvolatile products are formed that alter functional, sensory, and nutritional qualities of oils. During the past 30 years, scientists have reported extensively on the physical and chemical changes that occur during frying and on the wide variety of decomposition products formed in frying oils. This chapter will review the physical and chemical changes in oils during frying, including reactions that occur in the frying process. In addition to discussing the degradation products formed, their effects on oil stability and quality of fried food will be included. Methods to measure oil deterioration will be discussed in terms of their significance, advantages, and limitations.

II. PROCESS OF FRYING
A. Changes in Oils During Heating and Frying
1. Physical Changes
Deep-fat frying is a process of cooking and drying in hot oil with simultaneous heat and mass transfer. As heat is transferred from the oil to the food, water is evaporated from the food and oil is absorbed by the food [1] (Fig. 1). Many factors affect heat and mass transfer, including thermal and physical properties of the food and oil, shape and size of the food, and oil temperature [2]. Although this chapter primarily presents information about the chemistry of frying, the physics of frying is of interest.
in understanding how food fries, and how heated oils and their degradation products interact with fried foods. This knowledge will also help in optimizing the frying process by controlling the process to produce good-quality food and oil with longer fry life. Blumenthal [3] not only described the physical mechanisms that take place as a french-fried potato is fried, but also outlined procedures to optimize the frying process in order to increase frying oil life and decrease oil absorption into food. Blumenthal also proposed conducting basic frying studies using model foods that have surface-to-volume ratios characteristic of some fried food, including cotton balls (all interior volume, crispy exterior surface to represent battered and breaded chicken); french fried potatoes (significant interior volume, significant external surface), and potato chips (large surface area, little interior volume). Alexander et al. [4] compared corn, peanut oil, and partially hydrogenated soybean oils heated in open containers with those heated in a pressurized deep-fat frying system and reported that pressurized deep frying with fats resulted in less deterioration than open-vat heating, as shown by chemical analyses. In discussing physical changes in food during frying, Pokorny [5] reported that oil is absorbed into the fried material and adsorbed on its surface and that oil losses depend on the type of fried food; for example, potatoes absorb more oil than meat. Sun, Moreira, and Palau [2,6,7] have extensively studied the fundamental mechanisms of deep-fat frying using tortilla chips as model food and have suggested approaches to reducing chip fat content from a food engineering approach.

Physical and chemical changes in oils that occur during heating and frying are presented in Table 1. Although there are specific methodologies for quantitative measurement of degradation processes and products, some qualitative changes can also be determined subjectively by visual inspection. Although these practices are not recommended, many small-scale oil users, such as restaurants, discard frying oils when frying causes excessive foaming of oil, or when the oil tends to smoke exces-
Table 1  Effects of Physical and Chemical Reactions During Deep-Fat Frying

<table>
<thead>
<tr>
<th>Physical Changes</th>
<th>Chemical Changes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increased viscosity, color, and foaming</td>
<td>Increased free fatty acids, carbonyl compounds, high molecular weight products</td>
</tr>
<tr>
<td>Decreased smoke point</td>
<td>Decreased unsaturation, flavor quality, nutritive value (essential fatty acids)</td>
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</tbody>
</table>

sively, or when the oil color darkens. Other characteristics of abused oil include increased viscosity, off-odors such as acrid and burnt, and development of off-flavors in fried food. Of course, instrumental or chemical analyses are preferable to visual inspection in measuring frying oil deterioration, especially in research applications. The relationship of physical and chemical changes in oil is usually predictable since the many variables in the frying process affect the onset and magnitude of these changes, as will be discussed later in this chapter. Nonvolatile decomposition products eventually produce physical changes in frying oil, such as increases in viscosity, color, and foaming [8]. Chemical changes during frying increase the concentration of free fatty acids as well as carbonyl and polymeric compounds, and decrease fatty acid unsaturation. The effect of chemical changes on flavor quality of fried food and on the stability of the oil will be discussed in this chapter; however, potential toxic effects of degradation products on health will not be included because this topic is well covered in the literature [4,9].

2. Chemical Changes

Frying oils not only transfer heat to cook foods but also help to produce distinctive fried-food flavor and, unfortunately, undesirable off-flavors if deteriorated oil is used. During deep-fat frying various deteriorative chemical processes (e.g., hydrolysis, oxidation, and polymerization) take place, and oils decompose to form volatile products and nonvolatile monomeric and polymeric compounds (Fig. 1). With continued heating and frying, these compounds decompose further until breakdown products accumulate to levels that produce off-flavors and potentially toxic effects, rendering the oil unsuitable for frying. The amounts of these compounds that are formed and their chemical structures depend on many factors, including oil and food types, frying conditions, and oxygen availability [10]. In addition, these chemical reactions—hydrolysis, oxidation, and polymerization—are interrelated producing a complex mixture of products. The individual processes of hydrolysis, oxidation, and polymerization and their degradation products are described below.

a. Hydrolysis. As food is placed in oil at frying temperatures, air and water initiate a series of interrelated reactions. Water and steam hydrolyze triglycerides, which produces mono- and diglycerides, and eventually free fatty acids and glycerol (Fig. 2). Glycerol partially evaporates because it volatilizes above 150°C, and the reaction equilibrium is shifted in favor of other hydrolysis products [5]. The extent of hydrolysis is a function of various factors, such as oil temperature, interface area between the oil and the aqueous phases, and amount of water and steam because water hydrolyzes oil more quickly than steam [5]. Free fatty acids and low molec-
ular weight acidic products arising from fat oxidation enhance the hydrolysis in the presence of steam during frying [5]. Hydrolysis products, like all oil degradation products, decrease the stability of frying oils and can be used to measure oil fry life, e.g., free fatty acids.

b. Oxidation. Oxygen, which is present in fresh oil and is introduced into the frying oil at the oil surface and by addition of food, activates a series of reactions involving formation of free radicals, hydroperoxides, and conjugated dienoic acids. The chemical reactions that occur during the oxidation process contribute to the formation of both volatile and nonvolatile decomposition products. For example, ethyl linoleate oxidation leads to the formation of conjugated hydroperoxides that can form noncycling long chain, products or they can cyclize and form peroxide polymers [8]. The oxidation mechanism in frying oils is similar to autoxidation at 25°C; however, the unstable primary oxidation products—hydroperoxides—decompose rapidly at 190°C into secondary oxidation products such as aldehydes and ketones (Fig. 3). Secondary oxidation products that are volatile significantly contribute to the odor of the oil and flavor of the fried food [11,12]. If the secondary oxidation products are unsaturated aldehydes, such as 2,4-decadienal, 2,4-nonadienal, 2,4-octadienal, 2-heptenal, or 2-octenal, they contribute to the characteristic fried flavor in oils that are not deteriorated and can be considered desirable [12]. However, saturated and unsaturated aldehydes, such as hexanal, heptanal, octanal, nonanal, and 2-decenal, have distinctive off-odors in olfactometry analyses of heated oil. Fruity and plastic are the predominate off-odors of heated high oleic oils and can be attributed primarily to heptanal, octanal, nonanal, and 2-decenal [11].

Analysis of primary oxidation products, such as hydroperoxides, at any point in the frying process provides little information because their formation and decom-
position fluctuate quickly and are not easily predicted. During frying, oils with polyunsaturated fatty acids, such as linoleic acid, have a distinct induction period of hydroperoxides followed by a rapid increase in peroxide values, then a rapid destruction of peroxides [8,13]. Measuring levels of polyunsaturated fatty acids, such as linoleic acid, can help determine extent of thermal oxidation. Wessels [10] reported that oxidative degradation produced oxidized triglycerides containing hydroperoxide, epoxy, hydroxy, and keto groups and dimeric fatty acids or dimeric triglycerides. Volatile degradation products are usually saturated and monounsaturated hydroxy, aldehydic, keto, and dicarboxylic acids; hydrocarbons; alcohols; aldehydes; ketones; and aromatic compounds [13].

c. Polymerization. Polymerization occurs during frying, producing a wide variety of chemical reactions that result in the formation of compounds with high molecular weight and polarity (Fig. 4). Polymers can form from free radicals or triglycerides by the Diels–Alder reaction. Cyclic fatty acids can form within one fatty acid; dimeric fatty acids can form between two fatty acids, either within or between triglycerides; and polymers with high molecular weight are obtained as these molecules continue to cross-link. As polymerized products increase in the frying oil, viscosity of the oil also increases.

B. Factors Affecting Oil Decomposition

Understanding the mechanism of thermal degradation of a frying oil is difficult because it is affected by many variables, such as unsaturation of fatty acids, oil temperature, oxygen absorption, metals in substrates and in the oil, and nature of the food [14]. A list of factors that affect the processes of hydrolysis, oxidation, and polymerization and eventually frying oil deterioration are presented in Table 2. Frying oil degradation can be managed and even inhibited by controlling these factors.
For example, to help inhibit frying oil deterioration, choose a fresh oil with good initial quality, no prior oxidation, low levels of polyunsaturated fatty acids, and low amounts of catalyzing metals. Additions of antioxidants and antifoam additives may also help maintain oil quality [15–19]. The type of food being fried affects the resulting composition of the frying oil as fatty acids are released from fat-containing foods, such as chicken, and their concentration in the frying oil increases with continued use. Fat from fish will also change the fatty acid composition of the oil and decrease the frying oil stability. Breaded and battered food can degrade frying oil more quickly than nonbreaded food and decrease stability as well. However, even foods such as potatoes degrade oil stability because of the increased aeration produced as the food is added to the frying oil. Food particles accumulating in the oil also deteriorate an oil quickly; therefore, filtering oils through adsorbents will help remove these particles along with other oxidation products to enhance oil fry life.

The extent of these degradation reactions can be limited by carefully managing frying conditions such as temperature and time, exposure of oil to oxygen, continuous

Table 2  Factors Affecting Frying Oil Decomposition

<table>
<thead>
<tr>
<th>Oil/food/additives</th>
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<td>Unsaturation of fatty acids</td>
<td>Oil temperature</td>
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<td>Type of oil</td>
<td>Frying time</td>
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<tr>
<td>Type of food</td>
<td>Aeration/oxygen absorption</td>
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<td>Metals in oil/food</td>
<td>Frying equipment</td>
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<td>Initial oil quality</td>
<td>Continuous or intermittent heating or frying</td>
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<tr>
<td>Degradation products in oil</td>
<td>Frying rate</td>
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<tr>
<td>Antioxidants</td>
<td>Heat transfer</td>
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<tr>
<td>Antifoam additives</td>
<td>Turnover rate; addition of makeup oil</td>
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<tr>
<td></td>
<td>Filtering of oil/fryer cleaning</td>
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</table>

Figure 4  Polymerization reactions in frying oils.
or intermittent frying, oil filtration, and turnover of oil. Frying protocols of intermittent or continuous frying affect fry life. Perkins and Van Akkeren [20] found that cottonseed oil intermittently heated for 62 hours had as much polar material as oil heated continuously for 166 hours. They suggested that this may be caused by increased amounts of fatty acyl peroxides which decompose upon repeated heating and cooling, causing further damage to the oil. Replenishing the fryer with fresh oil is commonly done in most frying operations; however, in the snack food industry where more makeup oil is added than in restaurant-style frying, a complete turnover of the oil in the first 8–12 hours of the frying cycle can be achieved [2]. Levels of the reaction products in frying oil can also be affected by absorption into the fried food and evaporation [21]. However, accumulation of degradation products in the frying medium and their eventual incorporation in fried foods is of primary concern when commercial or industrial frying operations are carried out under abusive conditions [9]. Fritsch [1] pointed out that combinations of these factors (Table 2) determine the rate at which the individual reaction takes place. For example, in one operation, the rate of hydrolysis may be twice that of the rate of oxidation, whereas in another operation with different conditions, the reverse may occur.

III. DECOMPOSITION PRODUCTS

During frying, oils degrade to form volatile and nonvolatile decomposition products. Foods fried in deteriorated oils may contain a significant amount of decomposition products to cause potential adverse effects to safety, flavor, flavor stability, color, and texture of the fried food. Although volatile compounds are primarily responsible for flavor—both positive and negative—thermal polymers do not affect flavor directly. Therefore, thermal polymers may exist in an edible product, but the conditions leading to their formation are not usually encountered in commercial practice. Chang and coworkers [22] isolated the nonvolatile fraction as a brownish, transparent viscous liquid, which is indicative of a considerable amount of decomposition products in the oil.

A. Volatile Decomposition Compounds

In deep-fat frying, as oil is continuously or intermittently heated in the presence of air, both thermal and oxidative decomposition of the oil occurs producing both volatile and nonvolatile decomposition products (Table 3). Selke et al. [23] identified volatile odor constituents and their precursors from heated soybean oil, using model triglycerides [pure triolein, mixture of triolein (25%)-tristearin, and a randomly esterified triglyceride of stearic and 25% oleic acids] heated at 192°C in air for 10 minutes. Each model system produced the same major compounds, identified as heptane, octane, heptanal, octanal, nonanal, 2-decenal, and 2-undecenal. These seven compounds were unique to the oxidation of the oleate fatty acid in each triglyceride sample. Later, Selke and coworkers [24] analyzed pure trilinolein and mixtures of trilinolein-tristearin, trilinolein-triolein, and trilinolein-triolein-tristearin heated to 192°C in air. Major volatiles included pentane, acrolein, pentanal, 1-pentanal, hexanal, 2- and/or 3 hexanal, 2-heptenal, 2-octenal, 2,4-decadienal, and 4,5-epoxide-2-enal.
Table 3  Volatile and Nonvolatile Decomposition Products from Frying Oil

<table>
<thead>
<tr>
<th>Nonvolatile</th>
<th>Volatile</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monoacylglycerols</td>
<td>Hydrocarbons</td>
</tr>
<tr>
<td>Diacylglycerols</td>
<td>Ketones</td>
</tr>
<tr>
<td>Oxidized triacylglycerols</td>
<td>Aldehydes</td>
</tr>
<tr>
<td>Triacylglycerol dimers</td>
<td>Alcohols</td>
</tr>
<tr>
<td>Triacylglycerol trimers</td>
<td>Esters</td>
</tr>
<tr>
<td>Triacylglycerol polymers</td>
<td>Lactones</td>
</tr>
<tr>
<td>Free fatty acids</td>
<td></td>
</tr>
</tbody>
</table>

**B. Nonvolatile Decomposition Compounds**

Nonvolatile degradation products in abused frying oils include polymeric triacylglycerols, oxidized triacylglycerol derivatives, cyclic substances, and breakdown products [13]. Polymeric triacylglycerols result from condensation of two or more triacylglycerol molecules to form polar and nonpolar high molecular weight compounds. The nonpolymerized part of the oil contains mainly unchanged triacylglycerols in combination with their oxidized derivatives. In addition, it contains mono- and diacylglycerols, partial glycerols containing chain scission products, triacylglycerol with cyclic and/or dimeric fatty acids, and any other nonvolatile products. Rojo and Perkins [21] classified the degradation products as polar and nonpolar polymeric fatty acid methyl esters and monomeric fatty acid methyl esters with unchanged, changed (oxidized, cyclized, isomerized, etc.), and fragmented fatty acid esters. Clark and Serbia acknowledged that large declines in iodine values are needed for a significant amount of these polymers to form [9]. This is not usually a problem in the potato chip industry because iodine values do not change much as a result of a high oil turnover rate. However, oil used in small-scale batch frying operations such as restaurants, where oil turnover is low, may be more deteriorated. Chang et al. [22] studied the nonvolatile decomposition products from pure trilinolein, triolein, and tristearin produced under simulated deep-fat frying conditions at 185°C for 74 hours. Chromatographic, chemical, and spectrometric analysis indicated the presence of dimers in all three triacylglycerol mixtures. Similarly, Christopoulou and Perkins [25] isolated and characterized dimers in heated soybean oil. Dimers and higher polymers isolated from heated cottonseed oil at 225°C in the presence of air contained moderate amounts of carbonyl and hydroxyl groups [26].

**IV. MEASUREMENT OF DECOMPOSITION PRODUCTS: SIGNIFICANCE, ADVANTAGES, LIMITATIONS**

The physical and chemical changes occurring in frying oils and the many compounds formed in deteriorated frying oil have been extensively reported. Although these compounds often are used to measure degradation, many of the existing methods are based on measuring nonspecific compounds that may or may not relate to oil degradation or fried-food quality. Therefore, it is not surprising that frying is often described as more of an art than a science. In fact, the frying industry is still searching
for the ultimate criteria to evaluate frying stability of oils and fried-food flavor quality and stability. White [27] reviewed existing analyses to measure formation of volatile and nonvolatile components in order to detect deterioration in frying oils including the standard methods of polar components, conjugated dienes, and fatty acids, as well as rapid analyses such as dielectric constant. Croon et al. [28] compared methods to evaluate 100 frying oil samples using four quick test methods (Foodoil Sensor—dielectric constant, RAU Test, Fritest, and spot test) and two laboratory methods (free fatty acids and chromatographic analysis of triglyceride dimers) with a standard column chromatographic determination of polar compounds. Foodoil Sensor (FOS) Model NI-20 (Northern Instruments Corp., Lino Lakes, MN, USA) measures dielectric constant in frying fat relative to fresh oil; RAU Test is a colorimetric test kit that contains redox indicators reacting with the total amount of oxidized compounds. Fritest (E Merck, Darmstadt) is a calorimetric test kit sensitive to carbonyl compounds and the spot test assays free fatty acids to indicate hydrolytic degradation and free fatty acid. The Foodoil Sensor correlated with polar compounds more than did the RAU Test, Fritest, and spot test. The amount of free fatty acids was found to be an unreliable indication of deteriorated frying fat.

From a practical point of view, Fritsch [1] noted that commercial and industrial frying oil operators want to know the answer to one primary question: When should frying oil be discarded? Since there are many variables that affect oil degradation (Table 2), a specific method may be ideal for one operation but completely useless in another. Fritsch stated that determination of the end point of a frying oil is dependent on good judgment and knowledge of the particular frying operation, as well as on the type of frying oil and the analytical measurements used [1]. Some of the methods used to measure degradation products in frying oil are listed in Table 4 and are discussed in the following section. Schemes for the chromatographic separation of volatile and nonvolatile products in frying oil appear in Figure 5. After column chromatography, both polar and nonpolar compounds can be detected by high-performance size exclusion chromatography (HPSEC). Volatile compounds can be collected by several techniques, including direct injection, static head space, dynamic or purge-and-trap head space, and solid phase microextraction and analyzed by capillary gas chromatography.

A. Nonvolatile Decomposition Products

Paradis and Nawar [29] reported that nonvolatile higher molecular weight compounds are reliable indicators of fat deterioration because their accumulation is steady and they are not volatile. As mentioned earlier, the formation and accumulation of nonvolatile compounds are responsible for physical changes in frying oil (Table 1) [8]. Most methods for assessing deterioration of frying fats are then based on these changes. Nonspecific methods for measuring nonvolatile compounds in deteriorated frying oil include free fatty acids [1], iodine value [30], non–urea adduct–forming esters [26], viscosity [32], and petroleum ether–insoluble oxidized fatty acids [33]. White acknowledged that none of these methods has proved to be a good measure of heat abuse [27]. Melton et al. [34] noted that nonvolatile decomposition products are a better measure of degradation of a frying oil than volatile products are and concluded that more research is needed to determine the total polar components levels at which different frying oils should be discarded and to relate those levels to fried-food quality for each oil type.
Smith et al. [35] evaluated 65 samples of partially hydrogenated soybean oil used for frying battered chicken and french fried potatoes in fast-service restaurants. Frying times correlated highly with increases in dielectric constant, polar materials, and free fatty acids. Oleic and linoleic acids increased in the shortenings with hours of use, whereas stearic acid decreased because of contamination with chicken fat. Collected samples that had been discarded before 100 hours of frying time had values of 4.0 for the Foodoil Sensor, 1% free fatty acids and 27% polar materials which have been suggested as end points for discarding frying oil. Perkins [8] measured nonvolatile decomposition products in cottonseed oil and tallow to show that polymers increased with increasing heating time, and that cottonseed oil was deteriorated more by intermittent heating and/or added water than by continuous heating and no water addition. Cuesta et al. [36] measured polar components by HPSEC to investigate the thermo-oxidative and hydrolytic changes in frying oil. These researchers were able to quantitate triacylglycerol polymers and dimers, oxidized triacylglycerols, diacylglycerols, and free fatty acids.

Christopoulou and Perkins [25] recommended model systems such as pure fatty acids and triglycerides oxidized under simulated deep-fat frying conditions to control the various factors (Table 2) affecting the thermal-oxidative reactions and to facilitate the structure elucidation of the decomposition products such as thermal and oxidative dimers. Arroyo et al. [14] reported a linear correlation of $r = 0.99$ between number of fryings and amount of decomposition products, including total polar compounds,
triacylglycerol polymers, and triacylglycerol dimers. Although diacylglycerol levels were significantly correlated \((r = 0.945)\), free fatty acids were not significantly correlated \((r = 0.27)\) with the number of fryings. Arroyo and coworkers also found that hydrolytic changes paralleled thermoxidative changes, as evidenced by high correlations between levels of triglyceride polymers and triglyceride dimers (thermooxidative process) and diglycerides (hydrolytic process) with the number of fryings [14]. Dobarganes et al. [37], who measured triglyceride species and polar compound level and distribution, found no significant differences in the frying oils and lipids extracted from fried food for either total polar compounds or polar compound distribution. Thus, the study results indicated no preferential adsorption of altered oil compounds on the fried potato surface. Billek et al. [33] compared four methods to assess frying oils and reported good correlations between results with gel permeation chromatography (GPC), liquid chromatography (LC) on a silica gel column, polar and nonpolar components column chromatography (CC) on silica gel and petroleum ether–insoluble oxidized fatty acids. However, they found that measuring petroleum ether–insoluble oxidized fatty acids was time consuming and inaccurate. The GPC method was able to determine dimeric and oligomeric triacylglycerols in frying oil irrespective of the presence of oxidized compounds, whereas the LC method indicated the total amount of polar and oxidized compounds. Separation of polar and nonpolar components by CC was simple and quick.

Wessels [10] reported that methods to analyze frying oils, including measurement of peroxide value, benzidine value, petroleum ether–insoluble oxidized fatty acids, acid value, smoke point, UV absorbance, refractive index, iodine value, viscosity, color, and fatty acid composition, were of limited significance. Abdel-Aal and Karara [38] measured changes occurring in corn oil during heating and during frying of potato chips and onion rings by refractive index, acid value, peroxide value, total carbonyls, benzidine value, and oil color (which all increased) and iodine value (which decreased). These changes were more pronounced in oil that was used intermittently rather than continuously. Furthermore, onion rings were more detrimental to the oil than potato chips, possibly because of the breading material that accumulated in the oil. These investigators observed significant differences in the physiochemical changes of the oil extracted from the fried foods and the frying oil.

### B. Volatile Decomposition Products

Since many of the volatile decomposition products volatilize during frying, it is difficult to get an accurate representation of oil deterioration by instrumental and chemical analyses of these compounds. Methods that measure volatile compounds directly or indirectly include peroxide value, gas chromatographic volatile compound analysis, and sensory analysis (Table 4).

#### 1. Peroxides

Fritsch [1] noted that peroxide value is not a good measure of heat abuse in frying oils because peroxides are unstable at frying temperature (Fig. 6). Usuki et al. [39] confirmed this observation with pan frying (thin-film heating) of soybean oil, which produced high peroxide values of 230°C. No thermostable peroxides were detected after the oil was fractionated by silicic acid column chromatography.
Table 4 Methods to Measure Decomposition Products in Frying Oil

<table>
<thead>
<tr>
<th>Nonvolatile compounds and related processes</th>
<th>Method ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iodine value</td>
<td>AOCS Cd 1-25/93; AOAC 28.023 (31)</td>
</tr>
<tr>
<td>Fatty acid composition</td>
<td>AOCS Ce 1-6293 (30)</td>
</tr>
<tr>
<td>Total polar compounds</td>
<td>AOCS Cd 20-91 (30)</td>
</tr>
<tr>
<td>High-performance size exclusion chromatography</td>
<td>(36)</td>
</tr>
<tr>
<td>Free fatty acids</td>
<td>AOCS Ca 5a-40/93 (30)</td>
</tr>
<tr>
<td>Dielectric constant</td>
<td>(1)</td>
</tr>
<tr>
<td>Non-urea-adduct-forming esters</td>
<td>(26)</td>
</tr>
<tr>
<td>Color</td>
<td>AOCS Td 3a-64/93 (30)</td>
</tr>
<tr>
<td>Viscosity</td>
<td>(32)</td>
</tr>
<tr>
<td>Smoke point</td>
<td>AOCS Cc 9a-48/93 (30)</td>
</tr>
<tr>
<td>Foam height</td>
<td>(32)</td>
</tr>
</tbody>
</table>

Volatile compounds and related processes

| Peroxide value                             | AOCS Cd 8-53 (30)                |
| Conjugated dienes                          | AOCS Ti 1a-64 (30)               |
| Volatile compounds                         | AOCS Cg 4-94 (30)                |
| Sensory analysis of odor and flavor        | (48)                             |

2. Volatile Compounds

The fatty acid composition of frying oils has a major effect on the volatile compounds detected in the oil and on the flavor of the fried food. Although frying oils are complex mixtures of triacylglycerols, a wide variety of fatty acids, and many minor constituents, degradation compounds are primarily from the fatty acids [5]. Chang et al. [22] found that 79 of 93 compounds identified in corn oil and 64 of

Figure 6 Changes in amounts of volatile and nonvolatile decomposition products during the frying process.
100 compounds identified in hydrogenated cottonseed oil were also detected in pure triolein and trilinolein after all were heated under simulated frying conditions. Even though the fatty acid composition of the two oils differed greatly, approximately half of the compounds were the same. As expected, unsaturated fatty acids contribute significantly more to the formation of volatile compounds than those from the more stable saturated fatty acids, such as palmitic and stearic.

Chang et al. [22] concluded that identifying volatile compounds in frying oil and fried food is important because these compounds help in understanding the chemical reactions that occur during frying and because flavor of deep-fat–fried food is partly attributable to the volatile compounds. These researchers identified 220 volatile compounds from corn oil, hydrogenated cottonseed oil, trilinolein, and triolein after simulated frying conditions. Macku and Shibamoto [40] collected volatile compounds formed in the head space from heated corn oil and identified 18 aldehydes, 15 heterocyclic compounds, 13 hydrocarbons, 11 ketones, 4 alcohols, 3 esters, and 7 miscellaneous compounds. Takeoka et al. [41] isolated and identified volatile constituents of unidentified frying oils by simultaneous distillation–extraction and fractionation by silica gel column chromatography as 1-pentanol, hexanal, furfural alcohol, (E)-2-heptanal, 5-methyl furfural, 1-octen-3-ol, octanal, 2-pentylfuran, (E)-2-octenal, nonanal, (E)-2-nonenal, and hexadecanoic acid. Chung et al. [42] used gas chromatography/mass spectrometry (GC-MS) to identify 99 volatile compounds in the head space of peanut oil heated to 50, 100, 150, or 200°C for 5 hours, including 42 hydrocarbons, 22 aldehydes, 11 fatty acids, 8 alcohols, 4 furans, 2 esters, and 2 lactones. Total amounts of all identified volatiles increased as oil temperature increased. Chang and coworkers [43] identified 53 volatile flavor compounds from potato chips, including 8 nitrogen compounds, 2 sulfur compounds, 14 hydrocarbons, 13 aldehydes, 2 ketones, 1 alcohol, 1 phenol, 3 esters, 1 ether, and 8 acids.

Twenty-six volatile compounds were identified and quantified from aged potato chips fried in partially hydrogenated canola oil or cottonseed oil [44]. The samples fried in cottonseed oil had higher concentrations of aldehydes but heterocyclic compounds levels were not different. No differences in peroxide values were found between oil type. Neff et al. [11] reported that 32 volatile compounds in triolein heated for up to 6 hours at 190°C had identifiable undesirable odors by olfactometry-GC-MS, whereas only 18 volatile compounds had identifiable undesirable odors in heated trilinolein. For the same heated oils, Warner et al. found that triolein had four volatile compounds with the desirable deep-fried odor but that trilinolein had seven compounds with the fried-food odor; however, more than 800 ppm of these compounds was found in trilinolein and only 30 ppm in triolein [12]. Care should be taken in interpreting data on volatile compounds in used frying oil because of the fluctuations in formation and degradation of the compounds at frying temperature (Fig. 6).

3. Sensory

Frying oil affects the flavor of fried food because these oils undergo chemical reactions and the reaction products contribute to the distinctive fried-food flavor [5,12] as well as to undesirable odors in deteriorated oils [11]. Therefore, flavor quality is affected by oil type, frying conditions, and degradation products. In addition, Gere [45] noted the positive relationship of initial freshness of frying oil and sensory properties of food fried in the oil. However, fried-food flavor is not optimal at the start of frying [46–49]. When oils are tasted before heating, they usually have little
flavor if properly processed. This low intensity of flavor continues during the early portions of the frying cycle because the typical fried flavor develops as heating and frying time increase. Food processors often heat oils or fry preliminary batches of food to condition the oil to develop this flavor. Some oils develop this characteristic deep fried-food flavor more quickly than others depending on the fatty acid composition of the oil. For example, Warner et al. [48] found in previous research that cottonseed oil with high (50–55%) linoleic acid produces significantly higher intensity of fried-food flavor in potato chips and french fried potatoes than do oils with low (10%) linoleic acid, such as high (80–90%) oleic oils. As the fatty acids decompose in high temperature conditions, the volatile degradation products produce characteristic flavors. Some oxidation products, such as 2,4-decadienal, are important in the formation of typical deep-fried flavor. Flavor improves after the first stage of frying and becomes less acceptable during the last stage. Frying conditions should be adjusted so that optimal flavor characteristics are maintained for as long as possible during the frying cycle.

Sensory evaluation is still the method most often used by different countries to determine when to discard a frying oil [34]. Billek [33] reported that scientific groups in Germany used sensory assessment of a used frying oil; however, if this method did not give a clear indication that the oil was deteriorated, then instrumental or chemical analysis was used to support a final decision on oil quality. More recently, recommendations from the Third International Symposium on Deep Fat Frying recommended that the principle quality index for deep-fat frying be sensory parameters of the fried food [50]. To further confirm oil abuse, total polar materials should be less than 24% and polymeric triglycerides less than 12% [50]. Sensory analysis of frying oil and fried-food quality may be conducted by analytical descriptive/discriminative panels using trained, experienced panelists [46–48,51] or by consumer panels using untrained judges [34]. Melton used consumer panels to find that the flavor likability of fried food is dependent on consumer perception and is affected by the type of oil used for frying [34]. In further studies, Melton and coworkers could find no differences in fresh chip flavor or likability scores between potato chips fried in partially hydrogenated canola oil or cottonseed oil when evaluated by a consumer sensory panel [44]. On the other hand, Warner found that a trained, experienced, analytical descriptive panel could detect differences ($P < 0.05$) in the type and intensity of flavors in fried food prepared in various oil types [48]. More research is needed to understand the relationship between fried-food flavor and the volatile and nonvolatile decomposition compounds produced in frying oils.

REFERENCES


I. INTRODUCTION

Processing seeds or animal tissues into edible oils can be broken into four sets of operations: recovery, refining, conversion, and stabilization. Oil recovery is often referred to as extraction or crushing when processing plant sources and rendering in the case of processing animal tissues. Oil extraction involves pressing the oil-bearing material to separate crude oil from the solids high in protein or washing flaked or modestly pressed material with solvent, almost always hexane. The defatted solids after pressing are known as cake and after solvent extraction as meal. The oil, crude oil because it contains undesirable components, such as pigments, phosphatides, free fatty acids, and off-flavors and off odors, must be refined to remove these contaminants and produce high quality edible oils. Refined oils consist primarily (>99%) of triglycerides and can be converted, usually by hydrogenation; but winterizing, fractional crystallization, and interesterification should also be considered conversion processes because they achieve different properties from the original oil, such as converting liquid oil into semisolid or solid fats. Plasticizing, tempering, and stehling are operations designed to stabilize crystal–oil mixtures used for shortenings and margarines.

II. OIL RECOVERY

For several thousand years, fats and oils have been recovered from oil-bearing seeds, fruits, and fatty animal tissues, and used for food, cosmetics, lubricants, and lighting fluids. Of the more than several hundred plants and animals that produce fats and
Table 1  Major Edible Fats and Oils in the United States and Methods of Processing

<table>
<thead>
<tr>
<th>Source</th>
<th>U.S. oil consumption* (million pounds)</th>
<th>Oil content (%)</th>
<th>Prevalent method of recovery</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soybean</td>
<td>15,655</td>
<td>19</td>
<td>Direct solvent extraction</td>
</tr>
<tr>
<td>Corn (germ)</td>
<td>1,397</td>
<td>40</td>
<td>Wet or dry milling and prepress solvent extraction</td>
</tr>
<tr>
<td>Tallow (edible tissue)</td>
<td>1,362</td>
<td>70–95</td>
<td>Wet or dry rendering</td>
</tr>
<tr>
<td>Canola</td>
<td>1,264</td>
<td>42</td>
<td>Prepress solvent extraction</td>
</tr>
<tr>
<td>Coconut (dried copra)</td>
<td>1,021</td>
<td>66</td>
<td>Hard pressing</td>
</tr>
<tr>
<td>Cottonseed</td>
<td>772</td>
<td>19</td>
<td>Hard pressing or prepressing or direct solvent extraction</td>
</tr>
<tr>
<td>Lard (edible tissue)</td>
<td>988</td>
<td>70–95</td>
<td>Wet or dry rendering</td>
</tr>
<tr>
<td>Palm</td>
<td>260</td>
<td>47</td>
<td>Hard pressing</td>
</tr>
<tr>
<td>Palm kernel</td>
<td>390</td>
<td>48</td>
<td>Hard pressing</td>
</tr>
<tr>
<td>Sunflower</td>
<td>320</td>
<td>40</td>
<td>Prepress solvent extraction</td>
</tr>
<tr>
<td>Peanut (shelled)</td>
<td>230</td>
<td>47</td>
<td>Hard pressing or prepress solvent extraction</td>
</tr>
</tbody>
</table>

*Data from Ref. 2.

Source: Ref. 1 except as otherwise noted.

oils in sufficient quantities to warrant processing into edible oils, only 11 sources are commercially significant in the United States (Table 1) [1,2]. In other countries the importance of each source may be different, and other sources not on this list may also be important, such as olive oil being consumed in large amounts in Mediterranean countries.

All oil recovery processes are designed to obtain oil triglycerides as free as possible from undesirable impurities; to obtain as high a yield as possible consistent with economics of the process; and to produce cake, meal, or flour (finely ground meal), usually high in protein content, of maximum value [3]. Three general types of processes are used to crush oilseeds: hard pressing, prepress solvent extraction, and direct solvent extraction. The extraction process of choice depends primarily upon the oil content of the source material, the amount of residual oil in the meal allowed, the amount of protein denaturation allowed, the amount of investment capital available, and local environmental laws concerning emissions of volatile organic compounds (VOCs).

A. Extraction of Oil from Oilseeds

The oldest oil recovery method is hard pressing, where the seed is pressed, usually after various pretreatments to enhance oil recovery, to squeeze the oil from the solids known as cake (Fig. 1). In the early years, lever presses and screw-operated presses, often driven by oxen or other work animals, were used; then, during the Industrial Revolution, batch hydraulic presses were introduced, which evolved at the turn of the century into continuous screw presses [4] connected to line shafts driven by steam engines. Today, continuous screw pressing and direct solvent extraction have
Figure 1  Depiction of hard screw pressing. (Diagram courtesy of Anderson International, Cleveland, OH.)
become the preferred processing methods because the oil is more completely recovered. For a long time, the rule of thumb has been that materials containing more than 30% oil require pressing, either hard pressing or prepressing prior to solvent extraction. Hard pressing involves squeezing as much oil as possible; prepress solvent extraction involves squeezing out only part of the oil before subjecting the partially deoiled material to more complete extraction with solvent. The recent adoption of the expander has largely done away with this rule, and even high-oil content materials can be solvent extracted today with no or little prior oil extraction [5,6]. Direct solvent extraction, without any prior pressing or expanding, has long been the most widely practiced method, the oil is more completely recovered, and the oil and meal are economically recovered undamaged by heat. Both prepress solvent extraction and direct solvent extraction are depicted in Figure 2.

1. Seed Storage

Oilseeds are often harvested at moisture contents higher than levels that allow for long-term storage and must be dried for safe storage. Increasingly, farmers are storing oilseeds, particularly soybeans, on the farm to take advantage of higher prices that are paid later in the crop year.

Storage for extended periods at moisture contents exceeding critical moisture levels will damage oilseeds, reducing the yields of oil and protein, and diminishing the quality of the oil (notably darker color and higher refining loss). Seeds at harvest are alive and respire, converting seed mass to CO₂ and other metabolites, albeit at low rates when the moisture content is below the critical moisture level. The critical moisture level for safe storage varies with the seed specie: usually, the higher the oil content, the lower the critical moisture value (Table 1). At moisture contents exceeding the critical moisture level, respiration rate increases, and the seed can even germinate and become subject to fungi attack. Respiration and germination liberate heat and, when there is insufficient aeration, the heat further accelerates these reactions. Under extreme conditions, the seed may become scorched or even catch on fire (especially cottonseed). Modern seed storage facilities employ temperature-monitoring systems to alert elevator and storage operators when seed temperatures exceed critical set points. Then the seed is moved to another bin to disperse hot spots and/or aerated (blowing air through the seed). The percentage of seed that is heat damaged is often a factor in the U.S. grades and standards (e.g., soybeans).

Overdrying can increase seed fragility, leading to excessive breakage during handling, storing, and processing. In the case of soybeans (a dicot), the cotyledon is prone to splitting into halves when the hull becomes separated from the cotyledon (meat) during conveying and transporting, a problem that becomes worse when the beans are overdried. Splits are undesirable because they are difficult to separate from foreign matter, and the oil deteriorates at a faster rate. Oil from soybean splits is higher in free fatty acids, phosphatides, iron, and peroxides due to activation of catabolic enzymes [7]. Oils from field- and storage-damaged seed usually are poor in flavor [8]. For these reasons, damaged kernels and splits are also factors in the U.S. grades and standards for soybeans.

When oilseeds are received at the crushing plant (Fig. 3), samples are often taken for analysis of moisture, foreign matter, damaged seed, oil, protein, and free fatty acid contents. Shipments arriving with similar values for these analyses may be segregated based on actions that are required to minimize further degradation.
Figure 2  Depiction of prepress solvent extraction and direct solvent extraction. (Courtesy of French Oil Machinery Co., Piqua, OH.)

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Figure 3  Oilseed mills. (A) Soybean mill in Iowa and (B) sunflower seed mill in North Dakota.
Seed containing more than the critical moisture content is immediately processed or dried. Seed with excessive foreign matter is transferred to scalping operations to reduce the level of contaminants. Foreign matter usually contains more moisture than the seed and the foreign matter tends to become concentrated at certain locations during placement into storage bins. Removing foreign matter reduces bulk moisture content and improves storability. Usually, foreign matter is also a factor in U.S. grading systems.

Both inside and outside storage systems are used (Fig. 4). Metal and concrete bins are used for inside storage. Cottonseed is often stored in special buildings having roof slopes the same as the angle of repose for the seed (Muskogee buildings). Some oilseeds are stored outside on concrete pads, and the piles may be covered with tarpaulins and/or equipped with air distribution systems.

Any handling, including normal harvesting, conveying, and transporting, increases seed damage. Nature has given seeds a high degree of subcellular organization that is affected by seed damage. Seed endosperm is composed of many cells containing oil and storage protein, which supply energy, nitrogen reserves, and other metabolites to support the growth of the embryo during germination. Storage protein (the greatest proportion of seed nitrogen) is concentrated in discrete bodies known as protein bodies; and lipids (predominantly triglycerides) are stored in spherozomes. Phospholipids are largely associated with pseudomembranes around protein bodies and spherozomes. Enzymes and cellular metabolites are present in the cytoplasm. Some of these enzymes, notably lipase, hydrolyze triglycerides increasing free fatty acid content, which must be removed by refining. Phospholipase activity may render some phospholipids non-hydratable and difficult to remove from the oil by means of normal refining procedures. Lipoxygenases, unique to soybeans and other legumes, oxidize linoleic and linolenic acids causing painty, green, beany flavors in protein; these enzymes may also reduce the oxidative stability of the oil.

In intact seeds, these enzymes are kept away from the oil by natural compartmentalization within the cells of the seeds. However, damage by weather, harvesting, and/or handling bruises the seed, breaking cell walls and membranes, allowing the oil and enzymes to come into contact with one another. This accelerates adverse reactions whose rates are dependent on temperature, moisture, and extent of damage.

2. Cleaning

The first step in processing oilseeds usually involves cleaning the seed (Fig. 5). Unless removed by cleaning operations, foreign matter reduces oil and protein contents and increases wear and damage to expensive processing equipment; in addition, foreign matter may adversely affect oil quality (especially color). Magnets are placed in chutes just ahead of processing equipment to prevent damage by tramp iron. Vibrating and/or shaker screens with or without aspiration are used to remove stems, pods, leaves, splits, broken grain, dirt, and extraneous seeds (Fig. 6).

3. Dehulling

Usually, it is desirable to remove the hull or seed coat that surrounds the oilseed meat. Hulls always contain much less oil and protein than do meats. Removing the hull reduces the amount of material that must be handled, extracted, and desolventized, thus increasing downstream plant capacity. The protein content of the meal is raised by removing the hull. Oftentimes, as with corn and sunflower seed, the hull
Figure 4  Cottonseed storage systems. (A) Uncovered, outside storage piles; (B) covered, outside storage piles; and (C) Muskogee buildings.
Figure 5 Steps in processing soybeans into meal and edible oil.

contains waxes on the outer surface as a natural protective mechanism of the seed. These waxes must be removed during latter oil-refining steps, often by winterization or cold centrifugation; otherwise, the wax may become insoluble at cold temperatures and make the oil unattractively cloudy and cause emulsions such as mayonnaise to separate. Removing the hulls can alleviate these wax problems. On the other hand, hulls can be helpful at times, such as by providing fiber to allow easier hard pressing or to enhance solvent drainage, and thus they are not always removed.

Dehulling must be done carefully, to ensure that the meat is not broken into too many small pieces, which would be difficult to separate from the hull. Also, crushing of meats during dehulling causes oil cells to be ruptured, freeing the oil. This damage should be minimized to prevent absorption of liberated oil by hulls that are removed. Either one of these problems increases the oil content of the hulls and
Figure 6  Seed cleaner (shaking screens and aspiration).

reduces the yield of oil. Some seeds, such as soybeans, are often dried, dried and conditioned, or heated prior to dehulling to help free the meat from the hull.

Corrugated roller mills or bar mills, which shear the seed, and impact mills, which shatter the brittle seed coat, are used. The equipment cuts or breaks the hull, respectively, to free the meat, a step often referred to as decortication. Sometimes, the combination of sizing the seed with shaker screens prior to decortication and using mills with different settings optimized for each seed size increases the efficiency of decortication. Hull/meat separation systems comprising one or more shaker screens, aspirators, and gravity tables are effective because the hull is often larger and almost always lower in density and more buoyant in an airstream than the oil-rich meat. Hulls may be blended back with meal to control protein level, sold as a separate coproduct for cattle roughage, or burned in boilers to generate steam and electricity in the process known as cogeneration.

When making edible flours, more complete removal of hulls is required. More than 90% of the hulls from soybeans must be removed to assure that the minimum
specification of 50% protein is met. Even in the livestock feed industry, the trend is toward meals containing higher protein and lower fiber contents. It is becoming increasingly difficult to produce soybean meal with high protein content because increasing farm yields are depressing protein content.

4. Hard Screw Pressing

Since flakes transfer heat more rapidly than do cracked meats, the seed is usually flaked [0.38–0.50 mm (0.015–0.020 in.) thickness] prior to hard pressing (Fig. 1), before cooking at 115°C over a 60-minute period. Initial stages of cooking should be done with moist heat (seed moisture maintained at 10%) injecting steam or spraying water into the top deck. Cooked flakes are dried in lower trays and should exit the cooker at less than 2.5% moisture [9]. The cooked and dried flakes are then conveyed to screw presses. If proper cooking methods are employed, in conjunction with a well-maintained, modern, screw press, a residual oil content as low as 3–4% can be achieved. Many screw press plants are quite small and do not optimally, if at all, flake, cook, and dry before screw pressing; so as much as 6–10% oil often remains with the cake. Ultimately, hard pressing with more and more pressure is self-defeating because application of pressure causes capillaries to be reduced in volume, sheared, and eventually sealed by coagulation of protein. This places the lowest practical limit at about 3% residual oil.

A screw press (Fig. 7) is basically a continuous screw auger designed to accept feed material and subject it to gradually increasing pressure as it is conveyed through the barrel cage. The barrel is composed of bars surrounding the screw and oriented parallel to the screw axis. The bars are separated by spacers decreasing in size toward the solids discharge end, which allow the oil to drain. A plug of compressed oil-lean solids, the cake, forms at the discharge end. Increasing pressure down the length of the barrel is achieved by increasing the root diameter of the screw, decreasing the pitch of the screw flights, and controlling the opening for the discharging cake by means of a choke. This design causes fresh material to be rammed against the plug [10,11].

Screw presses are composed of three sections: feeding, ramming, and plugging. The meats or flakes fall into a rapidly rotating feed screw, which feeds them into the pressing cage to expel entrapped air and squeeze out the easily removed oil. In some screw presses, the feeding section may be a separate vertical screw, or it may be mounted directly to the ramming screw. In either case, the feeding screw turns faster than the ramming screw. Knife bars with projecting nibs are clamped between the half cages to prevent cake slippage and rotation. Maximum pressure is developed in the ramming section as partially deoiled cake is rammed against the deoiled plug. The plug section provides the resistance against which the ramming occurs. The friction in the barrel generates heat that must be removed to achieve low residual oil. Some screw presses recycle cooled pressed oil over the cage to remove excess heat, while others use water-cooled shafts and bar cages.

In hard pressing, screw presses are choked to put maximum pressure on flaked, cooked seed while maintaining cake discharge. The oil drains from the screw press cage and is pumped or flows by gravity to a basin to allow settling of cellular debris (foots) that was removed with the oil. The cake is then ground into meal.

In the United States, hard pressing is largely limited to minor oilseeds (e.g., peanuts and rapeseed), or in areas where supplies are not sufficient for large-scale
solvent plants (in cottonseed). There is interest in hard screw pressing oilseeds to produce organically grown or certifiable non-generating modified vegetable oil or to comply with local laws that prevent construction of new solvent plants such as in California. A small amount of soybean is hard-screw-pressed where the meal has particularly high value (e.g., high rumen bypass meal for dairy cattle feed). Screw pressing of copra, peanuts, sesame, and cocoa butter also remains popular in developing countries where investment capital is limited.

5. Prepress Solvent Extraction

In prepress solvent extraction (Fig. 2), part of the oil that is easily removed, is pressed out as described above. The press is choked such that less pressure is developed. Consequently, less oil is extracted and throughput is increased. Usually, the oil con-
tent of prepress cake is 15–18%, and the partially deoiled cake is then extracted with solvent. The cake may be broken into pieces and even flaked to increase bulk density and extractor capacity, and to speed extraction. The remaining steps are the same as for direct solvent extraction, described in the next section.

6. Direct Solvent Extraction

Cell walls are impermeable to oil and nearly so to extraction solvents. Consequently, cell walls and membranes must be distorted or ruptured to get the oil out, regardless of whether solvent extracting or screw pressing is done. This requirement often calls for reducing the size of seed particles. Small pieces also transfer heat and moisture more readily during conditioning or cooking. But excessive size reduction reduces mechanical distortion during flaking. Flaking mills distort larger particles more than smaller particles [12]. Of course, particles must be small enough to pass into the nip between rolls of the mill. Heating prior to flaking reduces oil viscosity, inactivates enzymes, coagulates protein, ruptures some cell walls and membranes, and makes the seed particle plastic for subsequent flaking or pressing. Proper plastic texture is necessary to produce thin, nonfragile flakes with minimal fines and maximal cell distortion. The flaked material must have tenacious, thin structure, with porosity that allows transport of the oil or miscella (solvent-oil mixture).

It has long been recognized that flake thickness influences the rate of oil extraction. Oilseeds are conditioned prior to flaking by using vertical stack cookers or rotary steam tube driers to heat the seed to 70–80°C over 20–30 minutes, while maintaining 10.5–11.5% moisture [13,14]. This treatment makes the meats soft and pliable enough to be flaked with smooth-surfaced roller mills to low thickness, 0.25 mm (0.010–0.012 in.) without producing excessive fines, which adversely affect other operations.

Extraction has been likened to cleaning paint from a brush [15]. As such, key to getting the brush clean is getting good contact and penetration of the solvent; then there must be enough clean solvent to dissolve the solute (the paint in this example), and enough time and heat to quickly dissolve more solute. However, oilseeds extraction does not follow the single mechanism of leaching as the brush example implies and others have often assumed. Instead, oilseeds extraction involves a combination of leaching, diffusion, and dialysis [12,16–19], which results in an ever-decreasing rate of extraction as the relative importance of each mechanism changes during the course of extraction [20]. For flakes, the larger proportion of readily extractable oil is derived from ruptured cells, especially near the surface. The transfer of oil from distorted interior cells probably is governed by capillary flow, and the rate of oil transfer is partly dependent on viscosity of the miscella. A portion of the slowly extracted oil is contained within intact undistorted cells and must be transferred by osmosis. This transfer is very slow [21]. Presumably, the process of extruding flaked meats, known as expanding, shifts the relative importance toward leaching because nearly all the cells are ruptured and the collet structure is quite porous.

Another portion of the slowly extracted oil relates to slowly soluble extractable materials, such as phosphatides, free fatty acids, nonsaponifiables, and pigments, which contribute to refining loss. The best quality oil, high in triglyceride content, is extracted first, while with more exhaustive extraction, poorer quality oil is extracted. Thus, at low residual oil levels, the proportions of free fatty acids and phos-
phatides extracted are greater, as is the refining loss. However, current industry practice is to strive for the most complete extraction possible. Typically, residual oil contents range 0.5–1.0%.

Flake thickness [22–25] and solvent temperature [25,26] have profound effects on extraction rate, and empirical relationships to extraction time have been observed [27]. While these factors are easy to control by adept operators, general lack of understanding and appreciation often exist in practice. The moisture content of the flakes is another factor affecting the rate of solvent extraction [28]. In most cases, 9–11% moisture is ideal. Hexane and water are immiscible, and higher moisture contents interfere with the penetration of hexane. Lower moisture levels reduce the structural strength of the flakes leading to the production of additional fines.

To reduce the amount of solvent used in extractors, countercurrent flow of the solvent to the flakes is used (Fig. 8) [29]. That is, the freshest flakes contact the oldest solvent and progress through the process until nearly oil-free flakes contact fresh solvent. Flakes enter the extractor through a plug vapor seal that allows the material to enter while keeping hexane vapors from escaping. The extractor is an enclosed vessel designed to wash, extract, and drain flakes.

Two principal types of extractor have been employed over the years: immersion extractors and percolation extractors. An immersion extractor immerses and soaks the material in solvent (an industrial example is the Hildebrandt U-tube extractor, and a laboratory example is the Soxhlet extractor). Generally, more solvent usage is required by immersion extractors. Few immersion extractors processing oilseeds remain; percolation extractors now dominate. In a percolation extractor, the solvent percolates by gravity through a bed of material (a laboratory example of a percolation extractor is the Goldfisch). The solvent flows over the surface of the particles and diffuses through the material during its downward circuitous travel. Miscella flows

![Figure 8](image-url)  
**Figure 8**  Flake flow relative to solvent flow. (Redrawn from Ref. 29.)
in successive passes through the bed, while the solvent spray and the bed move in opposite directions to each other. Percolation extractors also have larger extraction capacities in less space, and fewer operating problems are associated with these devices than with immersion extractors.

Most modern extractors are of the percolation type (Figs. 9, 10). The extraction principles employed by most extractors are the same, but there are different methods of achieving countercurrent flow of solvent to flakes. The shallow-bed chain extractor (Fig. 9) is one of today’s popular extractors and resembles a full-loop conveyor [30]. Flakes fed into an inlet hopper are conveyed down the first leg of the loop, where they are washed with moderately dilute miscella to extract surface oil and penetrate the cells. As the flake bed moves into the bottom horizontal section, full miscella is recycled through the bed for filtering, and then to a liquid cyclone for removing fines and, finally, to the evaporation system. Flakes are conveyed counterclockwise, through a progressively more dilute miscella washes, until a final wash with fresh solvent is used in the top horizontal section of the loop. The latter half of the top loop is used for drainage, after which solvent-laden spent flakes pass to the desolventizer. The following advantages are claimed: the shallow bed (usually about 1 m) promotes drainage and, thus, low solvent carryover to the meal desolventizer and uniform contact of the flakes with solvent, since the bed is turned over while going up the right vertical leg.

Today hexanes (a blend of about 60% n-hexane with other hexane isomers) are the solvent of choice, although many other solvents were used during development of solvent extraction technologies. Alternative extraction solvents were extensively reviewed [31,32], but no suitable alternative has been developed. The primary disadvantage of hexane is its flammability and a price structure and supply that are tied to petroleum prices.

Flakes are conveyed to the extractor, where they are extracted for 30–60 minutes. Generally, less than 1% residual oil in the extracted material is achieved, and the amount is lower for soybeans (about 0.5%). The miscella contains 22–30% oil, and the solvent is separated from the crude oil by distillation and stripping columns. The miscella is heated under vacuum to evaporate the solvent, which is usually done in two stages. The first-stage evaporator concentrates the oil to about 90% and uses reclaimed heat from heated solvent vapors from meal desolventization. Steam is used to heat the second-stage evaporator, where the oil is concentrated to >99%. Most of the remaining solvent is removed in a disc-and-doughnut stripping column where evaporation is promoted by means of heat, vacuum (450–500 mm Hg absolute pressure), and steam sparging. Crude oil leaves the stripping column with less than 0.15% moisture and hexane [33]. Trading specifications require the oil to have a flash point greater than 250°C, which is equivalent to no more than 800 ppm of hexane.

The marc generally contains 30–32% solvent holdup, which must be recovered and recycled. Heat must be used to evaporate the solvent holdup from the meal. Live steam is also injected to aid heat transfer and to provide moisture vapor to strip the solvent. Regardless of the type of extractor used, the extracted flakes (spent flakes) must be drained of the solvent held by the material. The solvent that will not drain is referred to as solvent holdup, the solvent-laden flakes are called marc. Solvent holdup should be minimized because this solvent must be removed by evaporation.
Figure 9   One type of commonly used extractor. (A) Schematic drawing and (B) an actual installation. (Photo courtesy of Crown Iron Works Co., Minneapolis, MN.)
Figure 10 Additional commonly used extractors. (A) Deep-bed rotary basket extractor. (Courtesy of French Oil Mill Machinery Co., Piqua, OH.) (B) Deep-bed chain extractor. (Courtesy of De Smet, Edegem, Belgium.)
using heat. Greater solvent holdup increases the energy required for desolventizing the meal.

Toasting is often needed for feed meals to efficiently denature trypsin inhibitors (protease inhibitors in soybeans affecting protein digestibility) and the enzyme urease (soybeans), bind gossypol to protein (cottonseed), and improve protein digestibility. Of course, none of these objectives can be achieved without considerable protein denaturation and the accompanying loss of water solubility by the protein. However, depending on the method used, meals with great differences in protein solubilities or dispersibilities can be produced.

The preponderance of meal is used for feed, where extensive heat treatment is necessary to maximize feed conversion efficiency by livestock. A conventional desolventizer/toaster (DT) (Fig. 11) is usually composed of about six stacked trays, all with indirect heating. The first two employ live steam injection through nozzles within the sweep arms to evaporate the majority of the solvent. Meal advances down through the trays, and a series of gates and floats control the levels in each tray. The lower four trays are essentially toasting/drying sections, where the meal is held at a minimum temperature of 100°C, and the meal is dried to a value suitable for dryers.

![Figure 11](image.png)  
**Figure 11**  Meal desolventizing/toasting equipment.
that follow the DT. Drying at normal DT conditions to less than 17% moisture is detrimental to available lysine. However, meal should not leave the DT at more than 22% moisture, for this would result in prohibitive drying energy requirements.

In recent years, the Schumacher-type desolventizer/toaster/dryer/cooler has become widely accepted. This device consists of four trays: the top tray is for prede-solventizing; the second for desolventizing-toasting with injection of steam through its perforated bottom (achieving countercurrent use of steam relative to solvent evaporation); the third for drying, with hot air blown through its perforated bottom; and the fourth tray is for cooling by blowing cold air through its perforated bottom.

The flash desolventizer and the vapor desolventizer (Fig. 12) were developed to reduce protein denaturation and produce highly soluble protein food ingredients (e.g., protein isolates) from soybeans [34]. Integrating these systems with cooking systems produces edible protein flours with a broad spectrum of protein dispersibility characteristics. The system includes a desolventizing tube, a flake separator, a circulating blower, and a vapor heater. These units are arranged in a closed loop in which hexane vapor is superheated under pressure and continuously circulated. Solvent-laden flakes from dehulled soybeans are fed into the system and conveyed by the high velocity circulating vapor stream. The turbulent superheated vapor flow (157–166°C) elevates the temperature of the flakes to 77–88°C, well above the boiling point of hexane (65°C), in less than 3 seconds. Because the flakes enter the flash desolventizer at low moisture for a very short period and no steam is injected into the vapor stream, little denaturation of protein occurs. As the flakes travel through the tube to the cyclone separator, the greatest portion of the entrained hexane is evaporated. At this point, if care is taken during conditioning, the protein dispersibility index (PDI) of the meal protein will be 2–5% of the untreated seed (native protein). The substantially desolventized flakes are removed from the system through a cyclone with a vapor-tight, rotary airlock and go to deodorizers.

Vapor desolventizing is similar to flash desolventizing in that superheated hexane vapor furnishes the required heat energy. Flakes are contacted with hot hexane vapor in a horizontal drum equipped with an agitator/conveyor. Flakes from either system usually enter a deodorizer to be stripped of hexane traces using only indirect heat. A slow moving agitator gently tumbles the spent flakes. The PDI may be further reduced by up to 10 percentage units. The final PDI is controlled at the flake stripper. Sparge steam may be used to minimize solvent loss and produce low PDI products (50–65% PDI). If only indirect steam is used, medium-range PDI products are produced (60–75% PDI). If the stripper is bypassed or operated without any heat or steam, highly dispersible products can be produced (75–90% PDI). However, as PDI increases, more hexane remains with the flakes as they exit the system, 0.5–1.2% hexane for high PDI products.

7. Meal Grinding

Desolventized meal is generally ground so that 95% passes a U.S. 10-mesh screen and a maximum of 3–6% passes through a U.S. 80-mesh screen. Meal for edible purposes is ground, sized, and sold as grits in a wide variety of sizes and as flour (<U.S. 100 mesh).

8. Oil and Meal Storage

Both oil and meal must be cooled before placing in storage because degradation reactions are accelerated at higher temperatures. Neither product should be stored
any longer than necessary. Oil degrades through oxidation, although crude oil is more stable than refined oil because crude oil contains natural antioxidants that are removed in refining steps. Preventing water from contacting the oil is also important to prevent hydrolysis, which increases refining losses. Thus, to extend storability of oil, protection against water, heat, and air is important.

Figure 12  Setups for (A) flash desolventizing and (B) vapor desolventizing.
B. Extraction of Oil-Bearing Fruits

Oil palms and olives are two examples of commercially important fruits providing important edible oils. Palm fruit can provide two distinctly different fats, one from the fleshy mesocarp and the other from the seed kernels. While olives and palms are processed slightly differently, space allows discussion of the palm oil production system only (Fig. 13).

Palm trees are perennials grown on plantations, particularly in Malaysia. It takes 3 years for the plantings to mature sufficiently to bear fruit, and they produce for about 25 years. Palm fruits are hand-harvested and immediately transported to the mill, where they are quickly sterilized. The fruits are processed within hours after harvesting because the oil immediately begins to degrade after harvesting. The fruits are sterilized by heating under steam pressure (145°C) for 1 hour to inactivate the

![Diagram of steps in processing palm fruits](image)

Figure 13 Steps in processing palm fruits.
enzyme lipase, which otherwise would quickly hydrolyze the oil and increase refining loss [35]. Sterilization also aids in stripping the fruit from bunch stalks and precondition the material for subsequent steps.

The sterilized fruits are stripped from bunch stalks with drum-type strippers. The material is then sent to a digester where it is reheated (95–100°C for 2 minutes) to loosen the pericarp from the nuts and to break the oil cells. The material is conveyed to continuous screw presses similar to, but not quite the same as, those used for oilseeds, to extract oil from the fruit flesh, but not the kernel. The liquid extract, press liquor, from the screw press contains about two-thirds oil, one-quarter water, and one-tenth solids. The press cake contains fruit flesh fiber and nuts. Water must be added to the press liquor to facilitate satisfactory settling of solids after screening, an operation referred to as clarification. Oil is skimmed off the top and passed to a centrifuge (clarifier) and then to a vacuum dryer. The crude oil is cooled and placed in storage.

The press cake is conveyed by means of a breaking conveyor to an aspirator (a vertical air column), where the nuts fall into a rotating polishing drum at the bottom, and the fruit fiber is blown to a cyclone, where it is separated from discharge air. The fiber is used to fuel the steam boiler. The nuts are conditioned by drying to loosen the kernels from the shell and cool the nuts to harden the shell. The nuts are cracked in an impact mill into two or more pieces. The shells are separated from the kernel with winnowing columns and by hydrocloning or clay bathing. All three of these operations separate shells from kernels based on density differences. The kernels are dried and screw-pressed or solvent-extracted to produce palm kernel oil and meal [36]. For every 10 tons of palm oil produced, 1 ton of palm kernel oil is produced.

C. Recovery of Animal Fats and Marine Oils

Animal fats and marine oils are recovered from fatty tissues by the cooking process known as rendering. Both edible and inedible fats are produced; the inedible tallow and grease being the majority in the United States and used as an energy source in livestock feeds. Raw materials include animal offal, bones and trimmings from meat processors, fish species unsuitable for marketing as fillets and other fish products (menhaden, pilchard, herring, etc.), and fish cannery wastes. Until recent years, edible tallow and lard were used for deep-fat frying in fast-food restaurants; but recent consumer concerns over cholesterol and saturated fats have reduced sales in these markets. Increasingly, larger proportions are being used in margarine and bakery shortenings. The defatted solid material is high in excellent quality protein that can be sold for use in livestock feeds as meat and bone meal (45–54% protein), meat meal (52–60% protein), poultry by-product meal (58–62% protein), and fish meal (60–65% protein). Fish meal commands high prices because it is especially valued in poultry diets and aquaculture diets.

Both wet rendering and dry rendering methods are used. Regardless of the process used, the material is conveyed upon receipt to a crusher or prebreaker to break the material into small pieces (2–5 cm). The broken material is conveyed to either batch or continuous cookers, where heating and grinding evaporate the moisture, break down the fat cells, and release the fat, action not too dissimilar from frying bacon.
1. Wet Rendering

Wet rendering is the older method and involves cooking the material (in the presence of water) by steam under pressure \([172–516 \text{ kPa (25–75 psi)}]\) for 90–150 minutes [37]. When the added water comes only from steam, the process is known as *steam rendering* and this process is used to produce *prime steam lard*. The water, denatured protein, and other solids settle to the bottom, while the fat, being less dense, floats on top of the liquid. Water, known as *stick water*, is drained off, and the remaining *tankage* goes to a press for fat removal.

The presses may be either hydraulic batch type or continuous screw presses similar to those used for processing oilseeds. The high-protein solids portion is known as *cracklings* and typically contains 6–10% residual fat. The cracklings are hammer-milled and screened, with oversized particles being recycled to the mill, thus producing *meal*. The fat discharged from the press must be centrifuged and/or filtered. Most fish, such as anchovy and menhaden, are processed by wet rendering.

2. Dry Rendering

In the newer and more efficient dry rendering process, the material is cooked in its own fat (115–120°C) in agitated, steam-jacketed vessels for 1.5–4 hours, until the moisture has evaporated [37]. No steam or water is added. The cooked material is then passed across a screen to allow the free fat to drain. The remaining tankage is sent to a press, and the remaining steps are the same as for wet rendering.

III. REFINING

A. Background

Consumers usually want bland-flavored or flavor-neutral, light-colored, and physically and oxidatively stable oils. Crude oils are not usually considered to be edible until numerous nonglyceride compounds have been removed through operations collectively known as *refining*. However, some oils, such as olive, tallow, and lard, have been consumed without refining. Undesirable components of crude oils include small amounts of protein or other solids, phosphatides, undesirable natural flavors and odors, free fatty acids, pigments, waxes, sulfur-containing compounds (canola and rapeseed), trace solvent residue, and water. However, not all nonglyceride compounds are deleterious. Tocopherols protect the oil against autoxidation and provide vitamin E activity, and \(\beta\)-carotene provides vitamin A activity. Other phenolic compounds, such as sesamol in sesame oil, act as natural antioxidants. Unfortunately, some of the refining operations are not perfectly selective and also remove some beneficial compounds along with the targeted undesirable ones.

There are two major types of refining: chemical and physical. The major steps involved in chemical refining include degumming, neutralizing, bleaching, and deodorizing (Fig. 14). Physical refining removes free fatty acids and flavors by distillation, to combine the steps of neutralization and deodorization into one operation. *RBD oil* refers to oil that has been alkali-refined, bleached, and deodorized or oil that has been physically refined.

B. Degumming

Degumming (Fig. 15) is a water-washing process to remove phosphatides. Unless removed, phosphatides can spontaneously hydrate from moisture in the air during
Figure 14  Steps in refining and converting edible oils.

storage or in the headspace. Degumming may be conducted either as a separate operation or simultaneously with neutralization. In the cases of oils rich in phosphatides, such as soybean and canola oils, degumming is usually a separate operation. Hydration makes phosphatides insoluble in the oil, and they precipitate, yielding an oil that is unattractive because of unsightly sludge or gums. Phosphatides can degrade and cause dark colors when the oil is heated as in the later deodorization step. All soybean oil in the export trade is degummed [38]. Phosphatides are also surfactants and, if present in frying oils, can cause dangerous foaming. When hot oil foams up and spills over the rim of a cooking vessel, it may burn the user; if it contacts a flame, it will catch fire. Phosphatides, also known as lecithin, are important food emulsifiers and, in the case of soybean oil, oftentimes become economical to recover.

The gums are rendered insoluble in oil by hydrating them with 1–3% water. As a general rule, the amount of water should be equivalent to the hydratable phosphatide content of the oil. If single-bleached lecithin is to be recovered, then hydrogen peroxide may be added to the water; if double-bleached lecithin is to be recovered, then benzoyl peroxide may also be added [39]. Alternatively, the degummed lecithin alone may be treated with hydrogen peroxide or both. The mixture is intensively mixed and then agitated for 30–60 minutes at 60–80°C in a slow mixing
Figure 15  Process flow sheet for degumming. (Redrawn from diagram provided by Delaval Separator Co., Sullivan Systems, Inc., Larkspur, CA.)
vessel (hydration tank) to allow the phosphatides to become fully hydrated and to coalesce. Hydration of the phosphatides is not instantaneous, and adequate time must be allowed. Higher temperatures solubilize more phosphatides, and lower temperatures increase oil viscosity; either one reduces the efficacy of degumming [40]. The gums, being more dense than oil, can be removed by settling or filtering; more often, however, they are centrifuged out. The wet degummed oil is either dried (as described later) or immediately neutralized. Usually, about 90% of the phosphatides are removed by this process. The gums typically contain 25% water and 75% oil-soluble substances (of which one-third is neutral oil).

There are both hydratable and nonhydratable phosphatides. Of the 1–3% phosphatides in soybean oil, 0.2–0.8% are generally regarded nonhydratable. The phosphatides are composed of phosphatidylcholine, phosphatidylinositol, phosphatidylethanolamine, and phosphatidic acid. The first two are always hydratable; but, the latter two can complex with divalent metal ions, rendering them nonhydratable. *Acid degumming* and *superdegumming* make more of the phosphatides hydratable. Nonhydratable phosphatides remain oil soluble. The nonhydratable phosphatides are believed to be calcium and magnesium salts of phosphatidylethanolamine and, especially, phosphatidic acid, that arise from the enzymatic action of phospholipases when the cellular structure of the seed is damaged [41]. Nonhydratable phosphatides are particularly problematic in soybean oil.

Acid degumming is an improvement over conventional degumming described above and has become the usual practice in the U.S. soybean industry. A small amount (0.05–0.2%) of concentrated phosphoric acid (75%) is added to warm oil (70°C) followed by stirring for 5–30 minutes and degumming as described in connection with conventional degumming. Longer mixing times are often substituted for lower reaction temperatures. Phosphoric acid is added to make the phosphatides more hydratable by binding calcium and magnesium ions before adding water. Phosphoric acid pretreatment also partially removes chlorophyll from the oil.

Phosphatide content varies widely in vegetable oils but is highest in crude soybean oil (Table 2) [42–46]. Soybean oil is the only oil that is regularly degummed. The use of expanders in preparing soybeans for extraction almost doubles the usual phosphatide content of soybean oil. Only about half the phosphatide content of soybeans is extracted with hexane when preparing soybeans by flaking alone. Sometimes, the degumming operation is conducted at the mill so that the gums may be added back to the meal. Gums contribute digestible energy to livestock. The available U.S. supply of soy lecithin is about twice the volume that can be economically sold. The gums for lecithin production are dried and may be further purified and/or bleached. Soybean lecithin is a mixture of about 40% phosphatides (16% phosphatidylcholine, 14% phosphatidylethanolamine, and 10% phosphatidylinositol), 35% oil, 17% phytoglycolipids, 7% carbohydrate, and 1% moisture [49].

Recently *superdegumming* processes have been developed in which more of the phosphatides are rendered hydratable. A strong solution of citric acid is added to warm oil (70°C), and the mixture is stirred and cooled to 25°C to precondition the gums. Then water is added with stirring for an additional 3 hours to hydrate the gums. This process causes the phosphatides to form liquid phospholipid crystals, which are easily removed during centrifugation.

Another variation of degumming, *dry degumming*, is occasionally applied to oils relatively low in phosphatide content, such as palm, coconut, and peanut oils.
Table 2  Properties of Some Crude and Refined, Bleached, Deodorized (RBD) Oils

<table>
<thead>
<tr>
<th>Properties</th>
<th>Soybean(^b)</th>
<th>Cottonseed(^c)</th>
<th>Canola(^d)</th>
<th>Palm(^e)</th>
<th>Sunflowerseed(^f)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Crude RBD</td>
<td>Crude RBD</td>
<td>Crude RBD</td>
<td>Crude RBD</td>
<td>Crude RBD</td>
</tr>
<tr>
<td>Triglycerides (%)</td>
<td>95–97 &gt;99</td>
<td>NA &gt;99</td>
<td>NA &gt;99</td>
<td>NA &gt;99</td>
<td>NA &gt;99</td>
</tr>
<tr>
<td>Phosphatides (%)</td>
<td>1.5–2.5 0.003–0.045</td>
<td>0.7–0.9 NA</td>
<td>2.7–3.5 NA</td>
<td>0.006–0.013 0.012</td>
<td>0.5–1.0 NA</td>
</tr>
<tr>
<td>Unsaponifiable matter (%)</td>
<td>1.6 0.3 NA NA</td>
<td>0.5–1.2 NA NA</td>
<td>NA NA</td>
<td>NA NA &lt;1.3 NA</td>
<td>NA NA</td>
</tr>
<tr>
<td>Plant sterols</td>
<td>0.33 0.13 NA NA</td>
<td>0.37 NA NA</td>
<td>NA NA</td>
<td>0.036–0.062 0.011–0.016</td>
<td>NA NA</td>
</tr>
<tr>
<td>Tocopherols</td>
<td>0.15–0.21 0.11–0.18</td>
<td>0.11 0.06 NA</td>
<td>0.06 NA</td>
<td>0.06–0.10 0.04–0.06</td>
<td>0.05 NA</td>
</tr>
<tr>
<td>Hydrocarbons (squalene)</td>
<td>0.014 0.01 NA NA</td>
<td>0.01 NA NA</td>
<td>NA NA</td>
<td>0.02–0.05 NA NA</td>
<td>NA NA</td>
</tr>
<tr>
<td>Free fatty acids (%)</td>
<td>0.3–0.7 &lt;0.05 0.9–37 &lt;0.05</td>
<td>0.4–1.0 &lt;0.05</td>
<td>2.0–5.0 &lt;0.10</td>
<td>0.8–2.4 &lt;0.05</td>
<td></td>
</tr>
<tr>
<td>Trace metals</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Iron (ppm)</td>
<td>1–3 0.1–0.3 NA NA</td>
<td>NA NA</td>
<td>1.5 &lt;0.1 5–10</td>
<td>0.12 NA NA</td>
<td></td>
</tr>
<tr>
<td>Copper (ppm)</td>
<td>0.03–0.05 0.02–0.06 NA NA</td>
<td>0.10 &lt;0.01 NA</td>
<td>0.05 0.05</td>
<td>NA NA</td>
<td></td>
</tr>
</tbody>
</table>

\(^{a}\)NA, data not available.  
\(^{b}\)From Ref. 42.  
\(^{c}\)From Ref. 43.  
\(^{d}\)From Refs. 44, 45.  
\(^{e}\)From Refs. 35, 46, 47.  
\(^{f}\)From Ref. 48.
The oil is treated with concentrated acid to agglomerate the gums. The gums are then separated from the oil by being adsorbed to bleaching earth during subsequent steps of bleaching and filtering.

C. Neutralization (Alkali Refining)

The term neutralization comes from neutralizing the natural acidity of the oil emanating from the presence of free fatty acids. Some use the term refining to refer to neutralization. Neutralization is the most important operation in refining edible oils (Fig. 16). An improperly neutralized oil will present problems in subsequent refining steps of bleaching and deodorizing, and in conversion operations of hydrogenation and interesterification.

Neutralization is achieved by reacting the free fatty acid with caustic soda (sodium hydroxide) to form soap referred to as soapstock. Saponification refers to reactions between glycerides and sodium hydroxide to also form soaps. Neutralization must be done correctly or some of the glycerides will be saponified, resulting in increased refining loss. The oil, low in acid value, is termed neutral oil. Removing the soapstock must also be carefully done to prevent high losses of entrained neutral oil, a second means of increasing refining loss.

Soapstock is a coproduct of refineries in that it can be acidulated with sulfuric acid to produce a salable product. Once reacidified, the fatty acids (95% fatty acids) will separate in settling basins as 35–40% free liquid, so-called acid oil, from an emulsified layer (high in phosphatides) and a water layer. Most acid oil is used as a high-energy ingredient in livestock feed, but when market prices are attractive, it is sold to fatty acid producers, who distill it to produce feedstocks for various oleochemicals (e.g., surfactants and detergents).

In the case of cottonseed oil, proper and timely neutralization is important to achieve adequate removal of gossypol and oil that is low in red color. For reasons that are not clear, gossypol is adsorbed onto soapstock particles even though gossypol is unsaponifiable.

The amount and strength of sodium hydroxide used depend on the amount of free fatty acids present in the oil. Nearly all oils other than soybean and rapeseed oils are simultaneously degummed and neutralization. Free fatty acids form water-soluble sodium soaps, and any phosphatides become hydrated and water insoluble. The amount of sodium hydroxide used is termed treat. The proper treat produces adequately refined oil with the lowest refining loss. Excessive treat can saponify triglycerides and reduce the yield of refined oil. The proper treat is determined by titrating the oil to determine the free fatty acid content and using industry tables, such as those published in the Official Methods and Recommended Practices of the American Oil Chemists’ Society [50].

Proper neutralization is dependent upon using the proper amount of sodium hydroxide, proper mixing, proper temperature, adequate contact time, and efficient separation. As in acid degumming, some oils are preconditioned with phosphoric acid. That is, prior to neutralizing the oil is treated with 0.02–0.5% phosphoric acid at 60–90°C for 15–30 minutes, making the phosphatides less soluble in the oil and more easily removed. The proper amount of caustic is proportionately metered into the warm oil stream with good mixing and sent to retention, or dwell mixers (5–10 min mixing time). The emulsion is then thermally shocked by heating to about 75°C
Figure 16 Process flow sheet for alkali refining. (Redrawn from diagram provided by Delaval Separator Co., Sullivan Systems, Inc., Larkspur, CA.)
to break out the soapstock. Soapstock is removed from the oil by using continuous, disk-type centrifuges. Refined oil is then washed with soft water (10–20%) at 90°C and recentrifuged to remove most of the soap. The remaining soap is removed during bleaching. The presence of excessive soap going into the bleaching operation can reduce the effectiveness in removing colors.

D. Miscella Refining (Neutralization)

Alkali refining or neutralizing in the presence of hexane is known as miscella refining. In the case of cottonseed, it is desirable to carry out alkali refining as quickly as possible after extraction (about 6 h) at the extraction plant; otherwise, gossypol may become fixed in the oil, hence unremovable [51]. Also, carrying out alkali refining in the presence of hexane reduces viscosity of the oil phase and increases the density difference between the oil phase and the water/soap phase. This improves the separation efficiency and reduces refining loss. Usually the oil content of the miscella is concentrated to 40–60% oil. The oil is mixed with sodium hydroxide with high-shear mixers, sometimes using high-pressure piston pumps with homogenizing valves. The mixture is heated to 65°C to melt the soapstock and then cooled to 45°C, and the aqueous and oil phases are separated by centrifuging. Water washing is not required in miscella refining. The neutralized oil miscella must then be evaporated and the oil stripped, dried, and cooled. Miscella refining produces oil with better color. The soapstock is usually added back to the meal by way of the desolventizer/toaster and contributes digestible energy to the meal.

The oil then is marketed as once-refined oil. Once-refined oil is re-refined as any other oil after arriving at the vegetable oil refinery, but much less caustic is required because most of the free fatty acid content has already been neutralized. Although any oil can be subjected to miscella refining, the additional capital investment in the safety features allowing centrifuges to work with hexane is justified only for cottonseed.

E. Drying

The water saturation level in edible fats and oils is about 0.8%; but oils should contain less than 0.3%. Vegetable oils must be dried before heating for prolonged periods to high temperatures as in hydrogenation and deodorization; otherwise, hydrolysis can occur, recreating free fatty acids. Drying is accomplished by spraying the hot oil (115°C) into a vacuum tower (15 mm Hg absolute vacuum). The moisture content of the degummed and neutralized oil is reduced to less than 0.1%. The gums for lecithin production are also vacuum dried in this manner to 0.5% moisture.

F. Bleaching

The primary purpose of bleaching (Fig. 17) is to improve oil color by removing pigments with neutral clays, activated earths, synthetic silicates, silica gel, and carbon black. Other benefits of bleaching are the breakdown of peroxides and cleanup of residual traces of soaps and phosphatides. The primary pigments of concern are those that give red-brown (carotenoids, xanthophyll, gossypol, etc.) or green colors (chlorophyll). The process is generally done under vacuum because the usual bleaching clays can catalyze oxidation in the presence of air (or oxygen). Adsorbent is mixed
Figure 17  Process flow sheet for vacuum bleaching. (Redrawn from diagram provided by Delaval Separator Co., Sullivan Systems, Inc., Larkspur, CA.)
with hot oil (80–110°C) for 15–30 minutes to form a slurry. Mixing enhances oil contact with the adsorbent. The pigments are adsorbed onto the surfaces of various clays or earths (some may be activated by treatment with acid), even sometimes activated carbon, and the solids are removed by filtration. Activated earths are made from certain bentonites, specifically montmorillonite. Acid activation is believed to be achieved by replacing aluminum ions in the clay structure [52] with hydrogen ions by treating with sulfuric acid; excess acid is removed with water, and the activated earth is dried and milled [38]. The hard-to-bleach oils are normally done so with acid-activated clays.

Bleaching power seems to be a function of the clay’s bound acidity, and clays with high total acidity and a reasonable level of acidity are preferred. If a bleaching clay is washed completely free of residual acid, bleaching power is greatly reduced. Clay from which water has been removed gives better results than clays containing adsorbed water; but if the earth is dried to less than 10% moisture, its internal structure will collapse, reducing surface area and thus adsorptive power. This partly explains why heating the oil/clay slurry is important, to remove water that is adsorbed in the clay lattices.

About 0.2–2% bleaching clay is usually used, the precise amount depending upon the amounts of pigments present. At the low end, 0.2–0.4% is used for soybean oil, while rice bran oil requires considerably more (3–5%). In addition to removing pigments and residual soap, bleaching takes out trace metals and some oxidation products. It is important to remove as completely as possible any residual soap (typically <10 ppm) because soaps can poison hydrogenation and interesterification catalysts, reducing their activities. Bleaching may be done batchwise or continuously.

Usually the earth is mixed with a small amount of the oil at cool temperatures (80°C), while the bulk of the oil is deaerated and heated to bleaching temperature (100–110°C). Time is not as critical as temperature, usually only 15–20 minutes are all that is required. Once the proper temperature and vacuum have been achieved, the oil/earth slurry is allowed to enter and become mixed with the bulk oil. Good mixing is important to allow for contacting the oil.

Filters are usually precoated with diatomaceous earth to enhance removal of the bleaching earth by leaf filters. Exhaustive removal of the earth is very important to oil stability because the earth acts as a pro-oxidant. Used bleaching medium is called spent earth. Spent bleaching earth contains some entrained oil, as much as 30–50%. The spent bleaching earth is blown with steam to reduce the oil content of the cake to about 20%. Some processors wash the cake with solvent to reduce the oil content to about 5%. Because of the large surface area that may be exposed to air and the catalytic effect of bleaching earth, spent bleaching earth is prone to spontaneous combustion and is regarded as a hazardous material by landfill operators. Disposal of spent bleaching earth is becoming a problem, and there is considerable interest in regenerating bleaching earth by extracting contaminants.

Edible oils should be pale yellow. Color is measured by the Lovibond tintometer, usually in red and yellow terms. Most finished edible oils are less than 10 yellow and 2.5 red, with high-grade shortenings being less than 1.0 red.

G. Dewaxing

Waxes can harm the appearance of bottled oil by causing unsightly cloudiness or sediments. Corn, rice bran, safflower, sesame, and sunflower seed oils are notorious
for problematic high wax contents (0.2–3.0%) and must undergo dewaxing; occasionally canola oil also has wax problems.

Waxes can be removed by cooling the oil to 6–8°C and filtering or centrifuging at cold temperatures, a process similar to winterization (described in more detail in Sec. IV.B) [52]. To get wax crystals large enough to ease separation, cooling must be done slowly over 4 hours, and the crystals should be allowed to mature for another 6 hours [40]. The oil is then carefully heated to 18°C and filtered.

Sometimes, dewaxing is accomplished simultaneously with removing the gums and/or soapstock by carrying out the centrifugation at cool temperatures. Sunflower seed oil is often predewaxed (from 1500 to 400 ppm) by cooling to 25°C for 24 hours and then degumming with a centrifuge at this temperature. Alternatively, sunflower seed oil may be simultaneously dewaxed and alkali-refined. After neutralization, as already described, the oil–soapstock mixture is cooled to 5–8°C and held there for 4–5 hours under gentle mixing; then the oil is mixed with 4–6% of water heated to 18°C. The soapy water phase wets and causes the small wax crystals to form a heavy suspension in soapy water. The soapy suspension is centrifuged to produce a wax/soapstock fraction and a refined, dewaxed oil [40]. Sometimes sodium lauryl sulfate is added to help wet the crystals [53].

H. Deodorization

The final step in refining fats and oils is deodorization (Figs. 18 and 19). Oils that are converted by a variety of processes are done so before deodorizing. The primary objective of deodorization is to remove compounds responsible for undesirable odors and flavors, such as residual free fatty acids (especially low molecular weight fatty acids), aldehydes, ketones, and alcohols. Deodorization also removes peroxide decomposition products; freshly deodorized oil should have a peroxide value of zero and free fatty acid content of <0.03%. These compounds are more volatile than are triglycerides and are preferentially removed. Lard is one fat that is often not deodorized, since the better grades have mild, unobjectionable flavors. Some losses of monoglycerides, sterols, sterol esters, tocopherols, and other natural antioxidants also result. The tocopherols are potent natural antioxidants and contribute significantly to the greater oxidative stability of crude oil compared to deodorized oil.

Deodorization is essentially steam distillation performed at high temperatures (180–270°C) and under high vacuum (3–8 mm Hg absolute pressure). Steam is sparged to carry away the volatiles and to provide agitation. Vacuum is usually provided by three to five stages of steam ejectors connected in series. Because deodorization is a mass transfer operation, deodorizers are designed to provide large surface areas and shallow oil depths. Only minor amounts of triglycerides are lost. The concentration of materials to be removed is in the range of 0.1–1% for most oils, and the original values should be reduced by over 99% [54]. The usual loss of oil weight is 0.2–0.8%. The amount of stripping steam ranges from 10–50 kg of steam per 100 kg of oil.

Factors that affect the efficacy of deodorization are the vapor pressures of the materials to be removed, the product flow rate, the intimacy of steam mixing with the oil, the absolute pressure achieved during deodorization, the temperature of deodorization (which controls the vapor pressure of the materials being removed), the sparge steam rate, and the time of deodorization [54,55].
Figure 18  Process flow sheet for deodorization. (Redrawn from diagram provided by Delaval Separator Co., Sullivan Systems, Inc., Larkspur, CA.)
Figure 19  Oil processing facilities. (A) Oil deodorizer and (B) margarine packaging plant in Melbourne, Australia.
Usually steam alone is not used to heat the oil because very high steam pressures would be required to heat the oil sufficiently. Rather, a eutectic mixture of diphenyl and diphenyl oxide, known by the trade name Dowtherm A, is used. This product has a boiling point of 258°C and at 304°C generates a pressure of only 110 kPa (16 psi).

Deodorization may be conducted in either batch, semibatch, or continuous vessels. The type of process largely depends on the volume and number of different products being processed. Batch deodorizers have cycle times of 6–8 hours. Continuous deodorizers are most suitable when a limited number of products are manufactured in very large volume. There are five stages in deodorization: deaeration, heating, deodorization/steam stripping, heat recovery/cooling, and final cooling. Stripping steam is provided through sparging rings and airlift pumps. After the deodorized oil has cooled, a small amount (0.005–0.01%) of citric acid is added to chelate metal cations so they would not promote oxidation and reduce shelf life. Crude oils usually have greater oxidative stability than refined oils. Indeed, many processors do not expose deodorized fats and oils to the atmosphere, but discharge oil into tanks blanketed with nitrogen and fill bottled oil under nitrogen.

Deodorization also removes any residual pesticide and hexane. Some pigments, such as β-carotene, are destroyed by the high heat in deodorization and, thus, the yellow and sometimes red colors are reduced. Deodorized oils have improved flavor, odor, and color. The high temperatures used in deodorization cause limited geometric isomerization. Although deodorization removes most peroxides, it cannot reclaim rancid oxidized oils.

Deodorizer distillate is condensed, recovered, and sold at higher prices per pound than the oil itself. Soybean deodorizer distillate typically contains 12.3% tocopherols and 21.9% sterols [49]. Deodorizer distillate may be further processed into valuable fractions rich in tocopherols (vitamin E), which are in high demand by the food and pharmaceutical industries. The sterols may also be purified and sold into the pharmaceutical industry for manufacturing various synthetic hormones.

I. Physical Refining

Physical refining (Fig. 20) is also known as steam refining. These terms are applied to the removal of the free fatty acids from the oil rather than reacting them with alkali, as well as to the removal of the compounds normally targeted by deodorization. Physical refining combines both neutralization and deodorization into one operation. Physical refining is always preceded by degumming and bleaching steps. The major advantage of physical refining is that the yield of oil is improved because there is none of the neutral oil loss that accompanies the production of soapstock. This process also affords the possibility of recovering fatty acids for the oleochemical industry without the need for acidulating soapstock and attendant wastewater production. The equipment used for physical refining is similar to deodorization, but with additional steam sparging trays.

Not all oils are suitable for physical refining. The oil must be low in phosphorus content (most comes from phosphatides). The exact upper limit for phosphorus in oil suitable for physical refining is a long-standing controversy, but it may be less than 5 ppm. Higher levels result in dark colored oils. Physical refining is also more appropriate for those higher in free fatty acid contents because the benefit of higher
Figure 20  Depiction of physical refining. (Redrawn from diagram provided by Delaval Separator Co., Sullivan Systems, Inc., Larkspur, CA.)

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yield justifies the cost of the process. Most regard soybean oil as being unsuitable for physical refining, or at best difficult to physically refine, while palm, rice bran, and coconut oils and animal fats are well suited to physical refining.

IV. CONVERSION

A. Background

Oftentimes, it is desirable or necessary to convert or transform highly unsaturated refined oils into more saturated forms; there may be the requirement for greater oxidative stability or altered physical forms of plastic and solid fats, or for less saturated forms to provide greater physical stability at cold temperatures (Fig. 14). The term conversion is often restricted to mean only hydrogenation, a process by which hydrogen is added to highly reactive, unsaturated, double carbon–carbon bonds. However, winterization, fractional crystallization, and interesterification, particularly directed interesterification, should also be regarded as modes of conversion, since these processes can also significantly change oxidative stability, physical stability, and functional properties of fats and oils, and their fractions. Stehling, votating, and tempering might also be considered to be conversion processes inasmuch as they are applied to the stabilization of fluid and plastic fats.

B. Winterization and Fractional Crystallization

The saturated fatty acids are not randomly distributed, and the oil can be fractionated into two or more fractions, differing in saturation level (as reflected by iodine value), and thus melting characteristics, oxidative stability, and functional properties. Both winterization and fractionation involve three stages: cooling the liquid oil to super saturation to form nuclei for crystallization, gradual cooling to remove latent heat of crystallization as the crystals grow in size, and separating the crystalline fraction from the liquid [56]. Supercooling to very low temperatures establishes excessive amounts of nuclei, which results in promotion of very small crystals that are difficult to recover by filtration.

1. Winterization

Winterization was first widely practiced on cottonseed oil. During the early years, the oil was stored in outdoor tanks, exposed to cold temperatures in the winter months. The oil would cloud, and as the crystals grew, they settled to the bottom of the tank. The crystals are composed of triglycerides containing more saturated fatty acids than the triglycerides composing the clear liquid oil. The clear oil was pumped off the lower, crystal-rich fraction, called stearin. Typically, 20–25% of the cottonseed oil comprised the crystallized stearin [43]. Cottonseed stearin was blended with other high melting fats to be used as shortening or margarine, where the high palmitic acid content contributes to improved crystal formation with improved functionality. Unwinterized cottonseed oil would cloud when stored under refrigerated temperatures and was unsuitable for salad oils and mayonnaise (the emulsion would break when crystals formed).

Today winterization consists of cooling bleached oil under refrigeration by passing the oil through continuous, chilling heat exchangers to cool the oil to
4–7°C, and then to tanks with slow agitation. The amount of agitation is critical: agitation is required to remove latent heat of crystallization (fusion), but too much agitation breaks up the crystals, making them more difficult to remove. Sometimes, compounds that act as crystal inhibitors are added to increase crystal size and aid filtration [57]. After several days to allow the crystals to grow, they are removed, usually with vacuum filters, but sometimes with plate-and-frame filters or centrifuges. The winterized oil (called winter salad oil) is then deodorized and packaged or bulk stored.

Some processors have carried out winterization in solvents, such as hexane. The solvent reduces viscosity, improves the efficiency of filtration, and increases yield of winterized oil. After filtration the solvent must be removed from the two fractions. This process is called miscella winterization.

In addition to cottonseed oil, canola oil is often winterized. Sunflower seed oil may also be winterized, not to remove saturated triglycerides, but to remove waxes that cause similar problems. Partially or lightly hydrogenated soybean oil that is winterized to remove saturated triglycerides (GSn) may be used for salad oils, which will have acceptable cold test values.

In a variation of winterization, some fats are pressed to achieve fractionation. Here a hydraulic press squeezes liquid oil from solid fat crystals. Hard butters for cocoa butter substitution in confectionery products and some specialty fats are produced by pressing of palm kernel and coconut oils [58].

2. Fractional Crystallization

So-called dry fractionation is a form of fractional crystallization very similar to winterization, but the term is usually reserved for more saturated fats (palm oil, palm kernel oil, animal fats), as opposed to liquid oils, where one or more cuts of crystallized fats, usually termed oleins (high in oleic acid), are removed from higher melting triglycerides, usually termed stearines (high in stearic acid). Temperatures higher than those normally reserved for winterization are used. For instance, crystallization of palm oil is carried out at 20°C to produce palm olein in about 70% yield and palm stearin (for shortening and margarine). A second fractionation at a lower temperature may be carried out to produce super palm olein (for frying and cooking oils) and palm mid-fraction (for cocoa butter substitutes). Fractional crystallization is used to make a number of cocoa butter substitutes from animal fats and from palm, palm kernel, and coconut oils.

Wet fractionation in various solvents (hexane, acetone, isopropanol, and 2-nitropropane) is also carried out on palm oil and hydrogenated soybean and cottonseed oils. Some confectionary fats and oils high in oxidative stability are produced by means of this form of fractional crystallization.

A third fractionation process is occasionally used, an aqueous detergent phase is mixed into preferentially wet, partially crystallized fat. The aqueous detergent phase contains 0.5% sodium lauryl sulfate, plus magnesium sulfate as an electrolyte. The crystals become suspended in the aqueous phase and are removed from the liquid oil by centrifugation. The water is removed from the crystals by heating the mixture and centrifuging. Both phases are washed with water to remove detergent and vacuum-dried to remove traces of water.
C. Hydrogenation

1. The Process

Hydrogenation was first used industrially to hydrogenate chemical feedstocks; it was first applied to whale oils in 1903 [59]. In 1909, hydrogenation was patented for use in producing shortening from cottonseed oil to replace lard. Hydrogenation is used for two purposes: to improve oxidative stability (by hydrogenating some of the double bonds to saturated ones) and to convert liquid oils or soft fats into plastic or hard fats (facilitating uses for which less saturated forms are unsatisfactory). Liquid oils are converted into shortenings and margarine fats; oils prone to rapid oxidation, such as soybean oil, are partially hydrogenated for use as salad and frying oils.

Unsaturated double bonds are converted to saturated bonds by addition of hydrogen (H$_2$). The reaction between the liquid oil and H$_2$ gas is accelerated by using a suitable solid catalyst; thus the reaction is heterogeneous involving three phases. Hydrogenation is exothermic, and heats about 1.7°C per unit drop in iodine value (IV) [60].

For successful hydrogenation, many of the crude oil impurities must be removed, and refining operations must be correctly carried out beforehand. Many of the contaminants (soaps, gums, sulfur, magnesium, potassium, chromium, zinc, and mercury) can poison the catalyst, reducing its activity. Canola and rapeseed oils are particularly notorious for high levels of natural sulfur content, which can cause problems in hydrogenation.

Both batch and continuous processes are used. For hydrogenation to occur, gaseous H$_2$, liquid oil, and solid catalyst must be brought together at a suitable temperature. Thus, hydrogenation is a mass transfer issue, and mass transfer of reactants is the rate-limiting factor (Fig. 21). H$_2$ is first dispersed as bubbles; then it must dissolve in the bulk oil, diffuse to the catalyst particle, and from there diffuse to the catalyst surface. Triglycerides also must diffuse to the catalyst surface, receive the H$_2$, and then diffuse out into the bulk oil. Therefore, a higher rate of reaction is achieved by raising the temperature and increasing agitation to disperse the gas in bubbles as small as possible. Small bubbles have more surface area for transfer per unit of mass for diffusion into the bulk oil. Agitation also keeps the film thicknesses small. Increasing gas flow rate helps to keep dissolved H$_2$ at high levels. The reaction is carried out in pressurized reactors because increased pressure increases the saturation concentration of H$_2$, hence the driving force for dissolution. Other factors that influence the reaction are the quantity and activity of the catalyst.

Pressurized reaction vessels are also used to contain the H$_2$, which is highly explosive should H$_2$, air (oxygen), and ignition come together. The reaction is normally conducted at 250–300°C over 40–60 minutes. The catalyst is mixed with a small part of the oil at room temperature. The bulk of the oil is pumped into the reactor, a vacuum is established, de-aerating the oil, and the oil is heated. Finally, the oil/catalyst slurry is added. The extent of reaction is followed by monitoring the refractive index (RI) of the oil, which is directly related to IV.

A catalyst increases the rate of reaction without being consumed in the reaction. Small amounts of catalyst are effective; in the case of hydrogenation, the usual amount is in the range of 0.01–0.02% of the weight of oil. Reduced nickel is the most widely used catalyst by the vegetable oils industry; but copper, platinum, palladium, and ruthenium are also effective. Nickel is the catalyst of choice because of
good activity, selectivity, filterability, reusability, and economical use. Copper has been occasionally used, but it must be completely removed to prevent accelerating oxidation. Since catalytic activity is a surface phenomenon, inducing pore formation increases surface area and thus activity [61]. However, the size and shape of pores in the catalyst are important; pores must be at least 10 nm in diameter in order to accommodate the transfer of triglycerides into them [60].

Catalyst promoters are substances that enhance the activities of catalysts without having catalytic activity themselves. It is believed that the promoter function is structural; that is, these substances somehow permit a larger number of active sites on the catalyst particle. For this reason, hydrogenation catalysts are commonly supported on siliceous materials, aluminum oxide, chromium oxide, cobalt oxide, or copper oxide.

Catalysts are expensive, but they can be reused numerous times. Hydrogenation catalysts can also catalyze oxidation. For these reasons, it is important to efficiently remove the catalyst from the hydrogenated oil and then recover it, which is accomplished by filtration. While surface area per unit weight increases as the size of the catalyst particle decreases, so does the relative difficulty of removing the catalyst by filtration. For that reason, the catalyst is often incorporated onto a support, such as 17–25% nickel fixed onto kieselguhr [60]. The catalyst is removed with leaf filters, and the filtered, hydrogenated oil is bleached to assure removal of all catalyst before deodorizing. This process, though termed post-bleaching, is carried out in the same manner as normal bleaching.

2. Selectivity

Selectivity refers to the relative rates of hydrogenation of specific fatty acids; selective hydrogenation is the opposite of random hydrogenation. Ideally, the most un-
saturated fatty acids are hydrogenated before the less unsaturated fatty acids have become hydrogenated (Fig. 22). High selectivity gives high oxidative stability for a given IV. For many applications one wants as much oxidative stability as is consistent with producing oil having no solid fat crystals at normal usage temperatures. Selectivity is always relative because perfect selectivity [i.e., all linolenic acid (18:3) converted to linoleic acid (18:2) before any linoleic acid is converted to oleic acid (18:1)] has not been achieved. However, producing catalysts with ever-increasing selectivity is a goal of catalyst suppliers.

3. Isomerization

Two types of isomerization spontaneously occur during hydrogenation: geometrical and positional. Only the extent to which isomerization occurs can be affected by processing conditions and catalyst selection. Geometrical isomerization refers to conversion of only cis double bonds to trans double bonds. Most unhydrogenated plant oils have only cis double bonds in their constituent fatty acids; however, milk fat and animal depot fats may have modest amounts of natural trans fatty acids, usually attributed biological hydrogenation by rumen bacteria.

Positional isomerization refers to the shift of double-bond position within the chain. If hydrogen atoms exist on the catalyst surface, the double bond opens, and hydrogen atoms add to the carbons atoms at either end [63]. When the catalyst has sufficient hydrogen atoms, another hydrogen atom is added such that the original double bond is converted to a saturated, single bond between carbons. When there are not sufficient hydrogen atoms to cover the catalyst, a hydrogen atom may be removed from either side of the partially saturated bond. This produces a new double bond.

![Figure 22](image)

**Figure 22** SFI properties of soybean oil hydrogenated under selective and nonselective conditions. (Redrawn from Ref. 53.)
bond, which may form in its original position or moved one carbon away, either up or down the chain. When the double bond re-forms, it may take either the \textit{cis} or the \textit{trans} configuration. These new double bonds may be half-hydrogenated and then dehydrogenated to re-form double bonds even further from the original position.

If hydrogenation is carried out at high pressure, low temperature, high agitation, high gassing rate, and low catalyst levels, then the catalyst is more saturated or covered with free hydrogen atoms, and there is little geometrical and positional isomerization. Also, under these conditions selectivity is low. However, under \textit{starved conditions} of limited hydrogen on the catalyst surface, when the reaction could proceed much faster if more hydrogen were present, both isomerization and selectivity are high.

Hydrogenation cannot be conducted without producing some geometrical and positional isomerization; but a wide range in the extent of isomerization is possible. The more times the catalyst is reused, the less selective it becomes, and the greater its tendency to produce isomerization. This has been attributed to natural poisoning of the catalyst by contaminants even in well-refined oil.

4. Effects on Physical and Functional Properties

Hydrogenation raises the melting point and reduces the IV of the triglycerides, usually to convert oils liquid at room temperature to semisolid plastic fats. Completely hydrogenated fats (IV < 1) are solid and brittle at room temperature and are used to add solids in baking shortenings. An advantage of hydrogenation is that a wide range of physical properties can be achieved, depending on how much H\textsubscript{2} is reacted with the oil. Thus, the processor and the food scientist have great flexibility in developing new products. Solid fat index (SFI) as determined by dilatometry and solid fat content (SFC) as determined by pulsed nuclear magnetic resonance (NMR) allow the analyst to describe the relative amounts of solid fat versus liquid oil over a temperature range usually reported at 10, 21.2, 26.7, 33.3, and 37.8°C, thus affording a measure of plasticity (especially important to margarines, baking shortenings, and confectionery fats).

Since \textit{trans} fatty acids melt at considerably higher temperatures than do \textit{cis} fatty acids, the former contribute considerably to melting and plastic properties of the fats. In some products, a high level of \textit{trans} fatty acids (as high as 55%) is desired for functional reasons, often to achieve proper melting characteristics (e.g., in cocoa butter substitutes). Margarines and shortenings usually contain substantial \textit{trans} fatty acids (12–33%) [44]. However, there has been and continues to be considerable debate about the effect of \textit{trans} fatty acids on health. Many now believe that \textit{trans} fatty acids are not as atherogenic as saturated fatty acids but not as healthy as unsaturated fatty acids [64].

Some catalysts are treated (e.g., H\textsubscript{2}S, SO\textsubscript{2}, CS\textsubscript{2}, and CO) to reduce sites available to hydrogen to increase \textit{trans} fatty acid production. There is also interest in developing catalysts that produce lower amounts of \textit{trans} isomers. Some of the semiprecious metals have this tendency (Fig. 23); but their costs are greater and recovery becomes even more important. Because some have higher catalytic activity than nickel, equivalent hydrogenation rates can be achieved at lower temperatures, reducing formation of \textit{trans} fatty acids, as well as energy usage [65]. Precious metals are normally supported on carbon. Once the catalytic activity is spent, the catalysts
are returned to the manufacturer for reclamation of the precious metal and remanufacturing of catalysts.

**D. Interesterification**

Interesterification, especially directed interesterification, can also be used to convert oils into more and/or less saturated fractions and ones with different functional properties from the original fat or oil or blend of two oils. Interesterification is described in much greater detail in Chapters 10 and 27.

**V. STABILIZING PHYSICAL FORMS**

Shortenings and margarines are *plastic*; that is, they have the appearance of solids in that they resist small stresses but yield to a deforming stress above a certain minimal value (the yield stress) to flow like liquid [66]. Shortenings and margarines are plastic at room temperature because they consist of two phases—solid fat crystals and liquid oil—and the solid crystals are sufficiently finely dispersed to be held together by internal cohesive forces (Fig. 24). Votating is conducted to produce the fine dispersion of solid crystals and entrained liquid oil. The dominant controlling factor for hardness is the relative proportion of solid crystals to liquid oil, followed by crystal size and polymorphic form (different crystal forms). The greater the proportion of crystals and the smaller the crystal size, the firmer the product, because there is greater opportunity for crystals to interassociate. The purpose of tempering is to facilitate transformation of the fat into the correct crystalline form.
A. Plasticizing

1. Plasticizing

Plasticizing is the most commonly used process for chilling and plasticizing margarines and shortenings. **Votator** is one manufacturer’s name for a swept-surface heat exchanger used to plasticize fat and thus the term **votation**. Molten fat is pumped into the first swept-surface heat exchanger known as an A unit, where the fat is supercooled to $15–25^\circ$C in $10–20$ seconds, depending on the product. Ammonia or **Freon** products are used as the heat exchange media in the exchanger jackets by means of a direct expansion refrigerant system. Oftentimes $5–25$ vol % of nitrogen, or in some cases air, is injected along with the melted fat and whipped or dispersed in the crystallized fat. The entrained gas makes shortenings white and may provide some physical leavening in bakery products.

The supercooled fat is then passed through one or more worker tubes, known as B units. Here the fat crystals are sheared while the heat of crystallization is dissipated. The temperature of the fat can rise $5–10^\circ$C as a result of the heat of crystallization. The fat at this point may be sent to a filling unit, or it may be subjected to a second but special heat exchanger known as a C unit. The final product will have a temperature of $13–24^\circ$C.

Firm table margarine is made with quiescent A units (lacking an internal agitator) replacing worker B units, whereas working B units are used to produce softer margarines.

2. Stehling

**Stehling** is a process that is often used to produce fluid shortenings, which are suspensions of fine crystals of solid fat in liquid oil. These products are pourable and pumpable, characteristics that are important to liquid frying shortenings, bread and cake shortenings, and nondairy products. Stehling is nothing more than stirring the oil/fat blend (and emulsifiers) after votating for a prolonged period at a precise temperature, to induce the formation of very small crystals that will remain dispersed in the liquid oil and not separate during reasonable transporting and storage periods.

B. Postprocessing Tempering

After plasticizing and packaging, shortenings are often moved to tempering rooms for $2–4$ days, where the temperature is controlled at $25–35^\circ$C, depending on the
product type and tempering time [67]. During tempering the fat transforms from one crystal form to another more stable form, a phenomenon known as polymorphism. Polymorphism refers to the property of fats to exist in several different crystalline forms, depending on the orientation of the molecules in crystallized fat. The crystals spontaneously transform from one crystal type to another, successively to the most stable form, higher in melting temperature. Both the rate and the extent of transformation depend on the fatty acid composition, the vortating conditions, and the temperature and duration of storage. Tempering expedites transformation to the most stable form. The least stable form is $\alpha$, and no fats are stable in this form. The next form is $\beta'$, and oils high in palmitic acid are stable in this form. This crystal form, characterized by smooth, small, fine crystals, is high functional in cake shortenings and margarines. The highest melting form is $\beta$, characterized by large, coarse, grainy crystals and is preferred in bread shortenings and confectionery fats.

Relative to untempered fats, tempering causes the fat to become slightly firmer above the tempering temperature and slightly softer at temperatures below the tempering temperature. The mechanism by which tempering occurs is not well understood, but it is believed to involve melting and recrystallization. Tempering serves to extend the plastic range of a shortening, rendering the product more functional. Untempered shortening may be grainy, brittle, and lacking in proper spreading qualities. Thus, tempering improves plasticity, creaming properties, and performance of baking margarines and shortenings.

VI. PROSPECTS FOR IMPROVED PROCESSES

Even though most regard the fats and oils industry to be mature, exciting new developments are in the process of adoption or testing. These developments involve advances in both processing techniques and equipment. In addition, new feedstocks modified through genetic engineering and traditional breeding could reduce or eliminate the need for some processing.

A. Expander Preparation

In recent years, expanders, an innovation pioneered in Brazil, have found acceptance for preparing flaked soybeans for extraction. Whereas with cottonseed, all the flakes are expanded, most soybean processors expand only part of the extractor feed, generally about 30%. As a result, part of the meal experiences more heat treatment than the unexpanded portion; but these differences are believed to have no practical significance. Expanding produces a porous collet for more rapid extraction, more complete drainage of solvent from the marc, higher miscella concentrations for lower evaporation costs, and higher bulk density for greater extractor capacity. Approximately 70% of the soybean mills in the United States now employ expanders.

In extrusion/pressing or expanding/pressing, whole soybeans are extruded or expanded, usually with an autogenous dry extruder (Fig. 25), followed by screw pressing. This nontraditional process is being used commercially in some unusual situations. In developing countries, for example, local processing of soybeans into crude cooking oils is attractive. Other applications of these extrusion/expansion techniques arise when specialty oilseeds are crushed for high-value oils, such as some genetically modified soybeans, and when high-value feed ingredients are produced.
(e.g., high-rumen bypass protein for dairy animals). The advantage is low capital investment, but a significant disadvantage is relatively low yield: about 6–7% residual oil is achieved compared with 0.5% residual oil for solvent extraction. Today, about 1–2% of the U.S. soybean crop is screw-pressed, but the percentage could increase if regulations on hexane emissions become more restrictive and mandated levels become more difficult to achieve.

B. Alternative Solvents for Extraction

The 1990 Clean Air Act is causing much concern over hexane emissions. n-Hexane, the main component of commercial hexane, is one of 189 Hazardous Air Pollutants listed in the Clean Air Act, and hexane will be regulated as both a criteria pollutant and a hazardous air pollutant. The emission limit as a criteria pollutant is 100 tons/year and as a hazardous air pollutant is 10 tons/year. To exceed either limit, a processor must obtain a federal operating permit, and there is an annual fee based on hexane consumption. Great strides have been made recently in reducing hexane loss, and today the average loss in a soybean plant would be about 0.2 gal/ton, down from 1.0 gal/ton 30 years ago. For a 2000 ton/day plant, however, this loss still translates into 460 tons of hexane per year.

While the industry continues to reduce hexane loss through engineering advances, a number of laboratories are researching alternative solvents (e.g., acetone, ethanol, isopropanol, isohexane, heptane) with less serious health and environmental risks. Enthusiasm for supercritical carbon dioxide has waned because of high capital costs and engineering problems in moving large quantities of solids through a reactor operating at very high pressure.
In a recent cottonseed plant trial, isohexane, a major component of commercial hexane, reportedly performed well and resulted in 38% steam savings [68]. But isohexane is even more expensive than hexane.

Work on ethanol and isopropanol has been going on for several years. The solubility of oil in these two solvents is temperature dependent, and this property can be used to advantage. High solvent-to-meal ratios are required because of low solubility compared with hexane. Acceptable energy usage is achieved by reducing evaporation costs by first chill separating. The full miscella is chilled to separate a heavier oil-rich phase containing more than 90% oil, which is stripped, and a lighter oil-lean phase that is recycled to the extractor. The extracted flakes are partially desolventized with mechanical presses followed by heat. Both solvent streams are used to wash the flakes. Although polar solvents usually extract poorer quality oil, the use of alcohols with chill separation produces good quality oil.

C. Membrane Filtration

Membrane technology to separate materials on the basis of molecular size has greatly improved in recent years [69]. Microfiltration, ultrafiltration, nanofiltration, and reversed osmosis are terms to designate different molecular weight separations. Membranes have been developed that are now stable to solvents and have greater selectivity. Membranes are being explored to concentrate oil in the miscella before evaporating.

Membrane filtration has considerable potential in degumming. When hydrated, the phosphatide molecule becomes oriented with the hydrophilic portion sequestered in the water droplet. In the nonaqueous environment of degumming, reverse micelles are formed. Micelles are large compared to the triglyceride molecules in which they are dispersed, and they are relatively easy to separate. The gums and pigments are concentrated in the 5% retentate, and high quality oil is recovered in the 95% permeate.

Membranes are also being explored to remove free fatty acids, and this approach appears to work. Perhaps more importantly, membrane-degummed oil is suitable for physical refining. Other applications for using membrane technologies in a vegetable oil refinery showing promise include miscella bleaching and hydrogenation catalyst removal.

D. Enzymatic Degumming

Both soybean and canola oils have high levels of nonhydratable phosphatides. Recently, enzymatic degumming for the conversion of nonhydratable phosphatides to hydratable forms has been perfected [70]. Water-degummed oil is treated with the enzyme phospholipase A2 after adjusting the pH to 5 with citric acid. Phospholipase A2 hydrolyzes the sn-2 fatty acid to form lysolecithin, which is easily hydrated and removed. The oil is sufficiently low in phosphatides to be suitable for physical refining.

E. Supercritical Fluid Refining

Supercritical fluids have been used to extract oil from oilseeds. The extracted oil is usually lower in phosphatides and free fatty acids. Although some report insufficient
selectivity, others claim success in using supercritical fluids to refine vegetable oils. Liquids are more suitable for supercritical fluid technologies than are solids in terms of material handling properties. Pumping against the high pressure is relatively easy.

F. Bleaching with Silica Gel

Bleaching clays are easily poisoned by residual soaps and phosphatides. Silica gel can be used to preserve adsorption capacity for pigments, especially problematic chlorophyll [71]. In the preferred method, the oil is contacted with silica gel before contact with clay occurs. When silica gel is incorporated, clay levels can be lowered by as much as 50%, reducing solid waste and neutral oil loss.

REFERENCES


I. CRYSTALLIZATION: GENERAL PRINCIPLES

Crystallization can be considered a subset of overall solidification. Solids are crystalline, semicrystalline, or amorphous. The crystallization process from solution first requires supersaturation; supercooling is a prerequisite for crystallizing from a melt. These phenomena lead to nucleation and crystal growth. This chapter emphasizes fat crystallization from the melt.

Once formed, crystals can have different shapes, called “habits” or morphologies. Stable crystals modify their habit, whereas metastable ones undergo phase transitions [1]. Both these processes result in polymorphic behavior, a behavior common to fats and other lipids. Further, most crystals “ripen” and disappear, as a result of changes in the degree of supersaturation [1]. Supersaturation “evolves” as crystal growth proceeds, and the liquid phase becomes less supersaturated. This reduced supersaturation results in a stability requirement for larger crystals, since crystals below a critical size will return to solution or the melt.

A. Supercooling

Crystallization requires a solute concentration greater than the concentration of the saturated solution. Observation of the schematic saturation–supersaturation curve in Figure 1 aids in understanding this phenomenon. The solid line represents a satu-
RATION or solubility curve. Below this, crystallization is impossible. Above this continuous line, the system is supersaturated: for example, at point 2, and crystallization is possible. Although crystallization is possible at point 2, it will not occur with agitation or seeding. This contrasts with point 3, where crystallization is spontaneous. The saturation curve results from thermodynamic factors; whereas the position of the unstable boundary (dashed line in Fig. 1) depends on kinetic factors, particularly the rate of cooling [2].

Review of the following equations, relative to melt systems, provides a conceptual understanding of how solubility (i.e., saturation) depends on the size of the solute. From Eq. (1),

$$\ln X = \frac{2\tau V}{rRT}$$  \hspace{1cm} (1)

where $X$ is the solubility increase versus crystals of infinite size, $\tau$ the surface tension, $V$ the molar volume, and $r$ the spherical crystal radius, it is clear that larger crystals would be less soluble. Equation (2),

$$\Delta T = \frac{2\tau VT}{r \Delta H}$$  \hspace{1cm} (2)

where $\Delta T$ is “supercooling” and $r$ is the smallest size of nuclei possible at a specific crystallization temperature $T - \Delta T$, indicates that the more a melt is supercooled, the smaller the critical radius necessary for stability.

B. Nucleation

Only when sufficient supercooling has been achieved can crystallization centers or nuclei form. A crystal nucleus is the smallest crystal that can exist in a solution with defined temperature and concentration [1]. Crystal nuclei were further distinguished from crystal “embryos” as molecular aggregates that continue to grow rather than redissolve [3].
Generally, it is accepted that both homogeneous and heterogeneous forms of nucleation occur. The former occurs in the absence of any foreign particle surfaces. The latter type occurs in practical systems when nuclei develop on the surface of solid impurities that are present. Chemical nucleation has also been referred to as a third, distinct type of nucleation [4]. Unlike heterogeneous nucleation, chemical nucleation results when organic agents that were added to a system dissolve and chemically react, activating the polymer. The ionic chain ends, which develop from polymer chain scission, aggregate and form nuclei. Therefore, it appears that chemical nucleation simply refers to chemical induction of heterogeneous nuclei rather than the more direct physical addition of solid heterogeneous nuclei.

Both primary and secondary nucleation have been described by many authors [1–3]. Secondary nucleation is relatively simple to understand. It results from fracture of growing crystals into smaller stable crystal nuclei. Nuclei stability is a function of nuclei solubility. This solubility, in turn, depends on the nuclei size at a given temperature. Nucleation theory addresses the concept of critical size regarding the development of stable primary nuclei.

An elegant discussion of these concepts was presented by Timms [2]. Two opposing actions exist when molecules attempt to aggregate. The first, energy evolution due to the heat of crystallization, favors crystallization as energy is released from the aggregating embryo. The second action, that of molecular surface enlargement, requires energy input to overcome surface tension or pressure. A stable nuclei will form only when the heat of crystallization is greater than the energy required to overcome surface energy. This relationship is represented in Eq. (3):

\[
\Delta G_{\text{embryo}} = 4 \pi r^2 \lambda - \frac{4 \pi r^3}{3 V_m} \Delta G_c
\]

where the first term on the right represents the surface energy contributions and the second depicts the volume contributions of the heat of fusion. Defining terms, \(\Delta G_{\text{embryo}}\) is the Gibbs free energy of the embryo, \(r\) is the radius of the nucleus, \(\lambda\) is the surface free energy per unit surface area, \(\Delta G_c\) is the molar free energy change resulting from the melt–solid phase change, and \(V_m\) is the molar volume [3]. The overall free energy will reach a maximum at some critical embryo size. Free energy will then tend toward a minimum in all embryos. This occurs through melting of smaller embryos or continued growth of embryos greater than the critical size.

C. Crystal Growth

Crystal growth continues as the properly configured crystallizing molecule diffuses to the proper place on the growing crystal surface. The rate of growth is directly proportional to supercooling and varies inversely with viscosity, since molecular diffusion is reduced as melt viscosity increases. Equation (2) predicts that small amounts of supercooling will lead to larger crystals versus greater supercooling leading to smaller crystals. These crystals will be relatively perfect, since the crystal attachment surface can become more precisely configured with the slower crystallization rate. Greater supercooling, by contrast, affords faster crystallization (provided the viscosity remains adequate for molecular diffusion) but also results in more crystal faults. Faults can arise as molecules from the melt attach to the crystal surface.
and have insufficient time to become optimally arranged before new attachments are made.

Boistelle [1] described one effect of the variable consequences resulting from different amounts of supercooling—the formation of “metastable phases.” These are simply thermodynamically unstable phases formed as a consequence of kinetic preference. In the case of crystallizing materials, the stable phase rarely forms first; rather, the free energy of the first crystal is closest to the free energy of the original melt. The kinetic preference for the unstable phase is a consequence of a lower difference in surface free energy [\( H \)] in Eq. (3)] between the melt and the crystal surface compared to the difference in the stable phase versus the melt. As with nucleation, the rate of growth is proportional to the degree of supersaturation. Since the solubility of a stable phase is less than that of a metastable phase, the nucleation and growth rates of the stable phase are expected to be faster. However, as discussed earlier, solubility curves are developed from thermodynamic data, whereas the initial nucleation and growth are governed by kinetics.

As crystallization continues, the degree of supersaturation in the system decreases. This then causes the critical crystal size to become greater. Therefore smaller crystals will dissolve and only larger crystals will grow. Eventually, only one crystal size will be stable. This process, called Ostwald ripening, occurs over a very long time possibly for years.

Sintering has been described as the formation of crystal bridges between preformed crystals in a semisolid dispersion of crystals and liquid [5]. In fats, the bridges are fat crystals having different thermodynamic and kinetic crystallization parameters relative to the precrystallized material. The bridge material can result from fractional crystallization and can consist of triacylglycerols or minor lipid compounds that fail to crystallize with the predominant lipids. Sintering has been measured in flocculation studies that assumed the adhesion between preformed crystals (sintering) resulted in greater flocculate volumes [5]. Sintering is important especially relative to the final texture and consistency of fat-based foods.

The process of sintering, like Ostwald ripening, occurs over a long time. As such it and Ostwald ripening are often called postcrystallization processes. Since, however, each describes crystallization phenomena, each is more precisely a crystal maturation process.

**D. Crystal Geometry**

Crystals are solids with atoms arranged in a periodic three-dimensional pattern [6]. Representation of the arrangement as a “point lattice” (Fig. 2) depicts each point having identical surroundings. All cells of a particular lattice are identical; therefore, any one cell’s dimensions and angles describe the lattice constants or lattice parameters of the unit cell (Fig. 3). The unit cell is the repeating unit of the whole structure. This compares to the concept of the subcell. As the name implies, a subcell is a smaller periodic structure within the real unit cell which defines the repeating unit of the whole structure [7]. This has particular relevance to long chain hydrocarbons, where the real unit cell is large and the subcell geometry is measured with X-ray diffraction. Only seven different cells are necessary to include all the possible point lattices. These correspond to the seven crystal systems into which all crystals can be classified (Table 1).
E. Crystal Polymorphism and Habit

Solids of the same composition that can exist in more than one form are referred to as polymorphic. The discussion of metastable versus stable in Sec. I.C alluded to this phenomenon. Crystal habit has simply been defined as the overall shape of the crystal [2]. From a crystallographic perspective, habit reflects the directional growth.
Table 1  The Seven Crystal Systems

<table>
<thead>
<tr>
<th>System</th>
<th>Angles and axial lengths</th>
</tr>
</thead>
<tbody>
<tr>
<td>Triclinic</td>
<td>All axes unequal and none at right angles</td>
</tr>
<tr>
<td></td>
<td>$a \neq b \neq c$ and $\alpha \neq \beta \neq \gamma$ and $\neq 90^\circ$</td>
</tr>
<tr>
<td>Orthorhombic</td>
<td>All axes unequal and all at right angles</td>
</tr>
<tr>
<td></td>
<td>$a \neq b \neq c$ and $\alpha = \beta = \gamma$ and $= 90^\circ$</td>
</tr>
<tr>
<td>Hexagonal</td>
<td>Two axes = 120° and the third at 90° relative to them</td>
</tr>
<tr>
<td></td>
<td>$a = b \neq c$ and $\alpha = \beta = 90^\circ$ and $\gamma = 120^\circ$</td>
</tr>
<tr>
<td>Cubic</td>
<td>All axes equal and all at right angles</td>
</tr>
<tr>
<td></td>
<td>$a = b = c$ and $\alpha = \beta = \gamma$ and $= 90^\circ$</td>
</tr>
<tr>
<td>Tetragonal</td>
<td>Two of three axes equal and all at right angles</td>
</tr>
<tr>
<td></td>
<td>$a = b \neq c$ and $\alpha = \beta = \gamma$ and $= 90^\circ$</td>
</tr>
<tr>
<td>Rhombohedral</td>
<td>All axes equal and none at right angles</td>
</tr>
<tr>
<td></td>
<td>$a = b = c$ and $\alpha = \beta = \gamma$ and $\neq 90^\circ$</td>
</tr>
<tr>
<td>Monoclinic</td>
<td>Three unequal axes having one pair not $= 90^\circ$</td>
</tr>
<tr>
<td></td>
<td>$a \neq b \neq c$ and $\alpha = \gamma = 90^\circ \neq \beta$</td>
</tr>
</tbody>
</table>

Source: Ref. 6.

within a crystal, whereas morphology describes the set of faces determined by means of the symmetry elements of the crystal [1]. This subtle distinction allows crystals of the same morphology to nevertheless occupy different habits.

Polymorphism is defined in terms of an ability to reveal different unit cell structures resulting from varied molecular packing. Polytypism refers to altered stacking direction of the crystal lamellae [8]. Each lamella is configured as identical polymorphs, but the direction of tilt of the hydrocarbon axis from the methyl end-group plane varies (Fig. 4).

II. FAT CRYSTALLIZATION

A. Lipid Classification

All fats are lipids, but the converse is not true. Therefore, a distinction must be drawn to provide an accurate perspective when considering fat crystallization as discussed in this chapter. Fats are defined as follows: “A glyceryl ester of higher fatty acids... Such esters are solids at room temperature and exhibit crystalline

![Image of polymorphism versus polytypism](image-url)

Figure 4  Polymorphism versus polytypism. (Reproduced with permission from Ref. 8.)
structure... The term ‘fat’ usually refers to triacylglycerols specifically, whereas ‘lipid’ is all-inclusive’ [9]. Included in the more general term “lipid” are hydrocarbons, steroids, soaps, detergents, all acylglycerols, phospholipids, gangliosides, and lipopolysaccharides [9]. Since all these lipids crystallize with many degrees of complexity, this chapter primarily addresses the crystallization of triacylglycerols, making only limited reference to other lipids.

Lipids have been classified based on their interaction with water [10]. Triacylglycerols belong to class I polar lipids. These are insoluble, nonswelling amphiphiles. Triacylglycerols will spread at the aqueous interface and form a stable monolayer. They have a low affinity for water compared to other class I polar lipids such as diacylglycerol and cholesterol. Class II polar lipids include many of the phospholipids, as well as glycolipids and monoacylglycerol. These, too, are insoluble; however, they swell because water is soluble in their polar moieties. A result of this interaction with water is the ability of class II polar lipids to undergo lyotropic mesomorphism and develop into liquid crystals. It has been suggested that mesomorphism of phospholipids influences the nucleation and solidification behavior of cocoa butter [11].

B. Triacylglycerol Crystal Packing Structure

Both the technical and biological functions of lipids are better understood with knowledge of their structural composition. X-ray analysis provides much of the structural information known regarding all lipids. Shipley [7] presented a brief history of the X-ray analysis of triacylglycerol single crystals. One of the first direct studies performed [12], the examination of the triclinic, \( \beta \) form of trilaurin, set the groundwork for determining the conformational nature of other, \( \beta \)-form triacylglycerols.

Of the seven crystal systems referred to in Sec. I.D, three predominate in the crystalline triacylglycerols [13]. Usually, the most stable form of triacylglycerols has a triclinic subcell with parallel hydrocarbon–chain planes (\( T \)). A second common subcell is orthorhombic with perpendicular chain phases (\( O \)). The third common subcell type is hexagonal (\( H \)) with no specific chain plane conformation [14]. Therefore the hexagonal form exhibits the lowest stability and has the highest Gibbs free energy, closest to the original melt. Figure 5 contains subcell representations of these three common triacylglycerol conformations.

Interpretation of X-ray crystallography data from trilaurin and tricaprin [15–17] resulted in representation of triacylglycerols in a tuning fork conformation when crystalline. The fatty acid esterified at the \( sn-1 \) and \( sn-2 \) positions of glycerol are extended and almost straight. The \( sn-3 \) ester projects 90° from \( sn-1 \) and \( sn-2 \), folds over at the carboxyl carbon, and aligns parallel to the \( sn-1 \) acyl ester. Molecules are packed in pairs, in a single layer arrangement, with the methyl groups and glycerol backbones in separate regions (Fig. 6).

The schematic view presented in Figure 6 represents simple, monoacid triacylglycerols. The polymorphic structures described for these simple triacylglycerols are valid for natural fats that contain complex triacylglycerols, given their common X-ray short spacings (axes normal to the chain direction) [13]. The structure depicted in Figure 6 illustrates a bilayer arrangement of the fatty acyl chains, which is the common packing structure for natural fats. However, this bilayer structure does not exist in all triacylglycerols [18–20]. Indeed, a trilayer structure has been demon-
This trilayer structure occurs when the sn-2 position of the triacylglycerol contains a fatty acid ester that is either cis-unsaturated or of a chain length different by four or more carbons from those on the sn-1 and sn-2 positions. Also, it was predicted that the trilayer structure would arise if the sn-2 position contained a saturated acyl ester with unsaturated moieties occupying the sn-1 and sn-3 positions [22]. Figure 7 depicts these varied layered structures.

When unsaturation results in a trans configuration around the carbon–carbon double bond, the crystal structure exhibits the normal bilayer appearance. The trans carbon–carbon double bond results in a linear chain configuration, unlike the bent chain configuration observed in the cis-unsaturated molecule. In fact, trielaidin (trans-C18:1) has the same polymorphic configuration as tristearin (C18:0) [13], although the phase transition temperatures of trielaidin are 30°C below that of tri-
stearin. Synthesized, stereospecific 1,2-dioleoyl-3-acyl-sn-glycerides with even carbon-saturated fatty acyl chains of 14–24 carbons have a trilayer packing structure [23].

III. POLYMERISM AND PHASE BEHAVIOR OF NATURAL FATS

As described earlier, solids with the same composition that exhibit different structural geometry are said to be polymorphic. The geometry of a particular polymorph confers unique physical properties beyond that of X-ray diffraction, although the X-ray diffraction pattern provides for unequivocal polymorph assignment [24]. Intramolecular and intermolecular conformation can also be inferred from infrared, Raman, and other forms of vibrational spectroscopy. Dilatometry and melting behavior are also used to evaluate the polymorphic nature of fats. Most multicomponent fats exhibit monotropic not enantiotropic polymorphism, and transformations proceed only from less stable to more stable forms [13].
Figure 7  Schematic comparison of triacylglycerol bilayer and trilayer structures: (a) tri-laurin (LLL), (b) 2-caproyldipalmitin (PCP), (c) 2-oleyldipalmitin (POP), and (d) 2-palmitoyldiolein (OPO). [Reproduced with permission from The Physical Chemistry of Lipids (D. M. Small, ed.), Plenum Press, New York, 1986.]

A. Nomenclature

The complicated melting behavior and polymorphism of fats resulted in confusion concerning terminology. Chapman [25] provided a good review of this controversy. It is now accepted that fats and triacylglycerols primarily occur in any of three basic polymorphic forms. The reference in Section II.B to the triclinic parallel (T), orthorhombic perpendicular (O), and hexagonal (H) subcells addressed polymorphic stability. The most stable and highest melting, T is the polymorph. Another polymorph, with variable stability and a melting point lower than , is . Phases with the hexagonal subcell have the lowest melting point and represent the polymorph.

X-ray diffraction provides not only the characteristic short spacing measurements, which define the lateral chain packing and subsequent polymorph assignment, but also measurements of the lamella layer thickness (the “ -spacing”). The -spacing depends on the length of the molecule and the angle of tilt between the chain axis and the basal lamellar plane. Although three basic polymorphs exist, subtle variations in -spacing lead to more than three polymorphic designations. Several of these were illustrated [26] and are shown in Figure 8. Figure 9 illustrates the orientation of the three primary polymorphs. The rarely isolated sub- form is very unstable and can be formed only at very low temperatures. The existence of more than one form results from chain tilt relative to the basal plane. This can be imagined by observation of the right-most projection of Figure 9 and the polymorph. A second polymorph would have a different interlamellar angle of tilt.
The $\beta'$ designation is maintained, since the subcell is still orthorhombic perpendicular ($O_{h}$); however a longer $d$-spacing results in a lower melting point and therefore a polymorphic designation $\beta'2$. As discussed earlier, this difference may also be referred to as polytypism.

Within groups having the same subcell, lower melting polymorphs are designated with a progressively higher subscript. The objective classification of acylglycerol polymorphs originally proposed [27] is also used for other lipids. This scheme (Table 2) refers largely to subcell packing and gives rise to subscripted forms having similar subcell dimensions. Finally, the bilayer or trilayer structure of triacylglycerol is designated with 2 or 3 following the polymorph description. For example, $\beta'_{2}$-2
designates a bilayer structure of a \( \beta' \) polymorph with the second highest melting point.

Application of the nomenclature to the most stable forms of various triacylglycerol systems, with \( sn \) positions designated A, B, and C for \( sn-1, 2, \) and 3, respectively, leads to the following:

**Table 2** Spectroscopic Parameters Used to Define Acyglycerol Polymorphs

<table>
<thead>
<tr>
<th>Polymorph</th>
<th>IR bands (cm(^{-1}))</th>
<th>X-ray short spacing (nm)</th>
<th>Subcell</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \alpha )</td>
<td>720</td>
<td>0.415 (st)*</td>
<td>Hexagonal</td>
</tr>
<tr>
<td>( \beta' )</td>
<td>719, 727</td>
<td>0.420, 0.380</td>
<td>( O_1 )</td>
</tr>
<tr>
<td>( \beta )</td>
<td>717</td>
<td>0.460 (st), 0.385 (st), 0.370 (st)</td>
<td>( T_1 )</td>
</tr>
</tbody>
</table>

*Strong line
Source: Ref. 13.
where A = B = C and all are saturated or trans-unsaturated, the form is \( \beta'-2 \);
where A = C and B is cis-unsaturated or has 4 or more carbons less than A and B, the form is \( \beta'-3 \);
where, in mixed saturated/unsaturated systems, A = B or B = C (asymmetrical), the form is \( \beta'-3 \);
where, in mixed saturated/unsaturated systems, A = C (symmetrical), the form is \( \beta'-3 \).

The packing requirements for \( \beta' \) are less stringent than for \( \beta \); therefore, mixed fatty acid triacylglycerols, such as those in lauric fats, tend to be \( \beta' \)-stable [28].

### B. Phase Behavior

Timms [24, 28] provided a complete review of the phase behavior of fats. A phase is a physical state of matter that is homogeneous and separated from other phases by a definite boundary. Therefore, if only the solid and liquid states of matter are considered, solid–solid, liquid–solid, and liquid–liquid phases can exist. Triacylglycerol mixtures are ideally miscible when liquid and therefore show no heat or volume changes when mixed. As a consequence, distinct liquid–liquid phases are not apparent. Many natural fats clearly exhibit several distinct solid phases with partial miscibility, which leads to compound crystals and solid solutions. Identification of these solid solutions with X-ray powder diffraction is effective, particularly with fats in equilibrium or relatively stable metastable forms. Thermal analysis using differential scanning calorimetry (DSC) provides for observation of the effect of changing temperature [28].

An idealized set of DSC thermograms is depicted in Figure 10. Melting profiles can be correlated to definitive X-ray determination, which allows subsequent estimation of polymorphic form using DSC alone. However, polymorph analysis using

![Figure 10](image_url)

**Figure 10** Schematic representation of polymorph identification by means of differential scanning calorimetry. [Reproduced with permission from *The Physical Chemistry of Lipids* (D. M. Small, ed.), Plenum Press, New York, 1986.]
DSC alone depends on thermal history, and other techniques may be necessary to sort out the complexity [22].

C. Milk Fat

At least 168 different triacylglycerol species have been identified in milk fat [29]. This complexity results in three distinct endotherms on heating of samples held at 26°C. As with many natural fats, the temperature at which crystallization occurs influences milk fat firmness, crystalline conformation, and percentage of solid fat [30]. Shear effects on milk fat crystallization have also been investigated [31]. Shear rate was directly related to crystal growth rate at crystallization temperature of 15°C and 20°C. At 30°C, shear was negatively related to growth rate, indicating the possibility of different growth mechanisms. However, the lower amount of supercooling at 30°C necessitates a larger crystal diameter for stability. Therefore, shear may simply interrupt the crystal growth, preventing development of the critical crystal size.

Fats may be thermally treated with a process referred to as tempering to produce specific attributes in a finished product. Tempering normally results in the formation of stable crystals having the proper size and in the proper amount [32,33]. Hardness variability in milk fat results from different thermal treatments. Less solid fat results when dairy butter is cooled slowly [34]. This is better understood considering the presence of three milk fat fractions with largely independent solid solutions [35]. A DSC thermogram (Fig. 11) depicts these independent fractions. The fractions are neatly defined as high-, middle-, and low-melting fractions (HMF, MMF, LMF, respectively). LMF is liquid at ambient temperature. Stable polymorphs of MMF and HMF were found to be a mixture of $\beta'$-2 + $\beta'$-3, and $\beta$-2, respectively. The role of LMF in the polymorphic transformations in milk fat fractions was investigated [36].

Figure 11  Differential scanning calorimetry melting curve of milk fat showing independent solid solutions.
LMF was found to facilitate the transformation of HMF to the $\beta$-2 form but has no effect on MMF, which stayed in the $\beta'$ form.

The various milk fat fractions and their different stable forms provide opportunities regarding applications to food and nonfood systems. Anhydrous milk fat fractions were evaluated regarding their effect on chocolate tempering [37]. Chocolate with HMF addition required normal chocolate tempering temperatures (<30°C) to produce adequate temper for molding, even though a high-melting endotherm existed after tempering at 31.1°C. This high-melting endotherm was attributed to the high-melting acylglycerols of HMF, which crystallized separately from cocoa butter. The incompatibility of the HMF with cocoa butter reported by the authors is not surprising, given the $\beta$-2 stable conformation for HMF and a $\beta$-3 stable form for cocoa butter.

D. Palm Oil

Palm oil is expressed from the pulp of the oil palm (Elaeis guineensis) fruit. It contains about 12 major triacylglycerols and is unique among vegetable oils because of the large percentage (10–15%) of saturated acyl esters at the sn-2 position. The free fatty acid composition is almost 5%, and there is a positive relationship between percentage of free fatty acid and hardness. At room temperature the oil appears as a slurry of crystals in oil. These phases persist even subsequent to tempering as measured by DSC [27]. The lower melting solid solution consists predominantly of 1-palmitoyl-2,3-dioleoyl-sn-glycerol (POO) and the high melting solution is dominated by 1,3-palmitoyl-2-oleoyl-sn-glycerol (POP). Three polymorphs were determined to be in palm oil [38]. These were $\beta_i$ (a sub-$\alpha$ form found only on rapid cooling to subzero temperatures), $\alpha$-2, and the stable $\beta'$ form. The stable $\beta'$ form is not surprising, given the heterogeneous triacylglycerol composition. The $\beta'$ stability has resulted in the addition of palm oil to oils destined for shortening or margarine, since $\beta$-tending fats can result in gritty textures. Small amounts of a palm oil $\beta$ form have been produced by means of thermocycling [39]. As expected, thermocycling of the stearin (high-melting) fraction led to almost 40% $\beta$ crystals after 36 days at 5°C.

E. Lauric Fats

Coconut oil and palm kernel oil are the two predominant lauric fats. The term “lauric fat” refers to the high percentage of lauric acyl esters in the triacylglycerol. The triacylglycerol composition of coconut oil is over 84% trisaturated, although at least 79 different triacylglycerol species have been identified. Less than 10% of the fatty acid composition is unsaturated [29,40]. The chain length of the acyl esters varies from C8 through C16. The polymorphic nature of coconut and palm kernel oil (PKO) as well as hydrogenated PKO and PKO stearin are similar and relatively simple. Two polymorphs have been identified: $\alpha$ and $\beta'$-2. The $\alpha$ form is fleeting and can be recognized only after rapid cooling, as it quickly transforms into the $\beta'$-2 polymorph [41–43]. This too is not surprising, given the mixed chain length and asymmetry of the triacylglycerol molecules. The melting point of these fats is sharp at 22°C for coconut and 25°C for PKO [29].

F. Liquid Oils

Evaluations of polymorphism in oils (fats that are liquid at room temperature) are limited. Cottonseed and peanut oils exhibit no $\alpha$ form and no $\beta$-2 form. These oils
simply crystallized in the sub-α form, sometimes referred to as β; because of its orthorhombic perpendicular subcell arrangement [43]. This form results only at very low temperatures. The sub-α form of cottonseed and peanut oils transformed into a stable β;2 form. Four other oils (corn, safflower, sunflower, and soybean) showed polymorphism similar to that of peanut and cottonseed, but these four fats developed a stable β-2 form.

G. Hydrogenated Fats

Complete hydrogenation eliminates the asymmetry, often leading to β’ stable polymorphs. Therefore, soybean, peanut, sunflower, corn, and sesame oils, having a large composition of C18 unsaturated fatty acids, are converted to hydrogenated fats having stearoyl esters and consequently show the stable β-2 form. More highly saturated fats, such as cottonseed, olive, palm, and cocoa, are converted to fats containing 1,3-dipalmitoyl-2-stearoyl-sn-glycerol (PStP), 1,3-distearoyl-2-palmitoyl-sn-glycerol (StPSt), or 1-palmitoyl-2,3-distearoyl-sn-glycerol (PStSt) [42,44]. The rearrangement of PStP into a stable β form is hindered by misalignment of the methyl end plane of the β’ unit cell [13], and a fat rich in this triacylglycerol will stay in the β’ form. Hydrogenated fats rich in StPSt can transform into a stable β form, since the interlamellar methyl end plane can rearrange more easily. The high PStSt fats have equally stable β’ and β forms, and any transformation to the β form occurs over a long period of time.

H. Cocoa Butter

Two reviews of the composition of cocoa butter provide a good understanding of the heterogeneity of this natural fat [45,46]. More than 98% of cocoa butter is simple lipid. More than 95% of this is triacylglycerol. Three triacylglycerols predominate: 1-palmitoyl-2-oleoyl-3-stearoyl-sn-glycerol (POS) composes about 40%, 1,3-distearoyl-2-oleoyl-sn-glycerol (SOS) makes up 27.5%, and 1,3-dipalmitoyl-2-oleoyl-sn-glycerol (POP) makes up about 15%. At least 15 minor triacylglycerols have also been identified. Free fatty acid values are reported to be 1.5%, and the concentration of mono- and diacylglycerol is about 2% of the simple lipid fraction. The concentration of phospholipid and glycolipids in cocoa butter varies by climate and method of analysis. These complex lipids constitute about 1% of original cocoa butter.

1. Cocoa Butter Polymorph Nomenclature

More is known about the phase behavior of cocoa butter than that of any other fat [28]. X-ray diffraction techniques have been used to define six polymorphs, each having a distinctive melting point [47]. Interestingly, refined cocoa butter was used in this work, and the possible effects of the compounds removed by refining appear to have been discounted. The existence of six polymorphs in a study with 12 different cocoa butters was confirmed [48]. Thermal analysis alone was used to evaluate the polymorphism of cocoa butter and mixtures of cocoa butter with other fats [49]. That study led to some of the confusion with terminology regarding cocoa butter polymorphs in that the previously defined [47] designations (from lowest, I, to highest, VI, melting point) were reversed regarding melting temperature. Also, six cocoa butter polymorphs were found in work combining microscopy and thermal analyses [50]. Table 3 lists the nomenclature and melting points of the cocoa butter poly-
morphs as determined by several groups. The definitions of cocoa butter polymorphs follow several conventions [27,47,51]. Consistency throughout this chapter requires the Wille and Lutton “I through VI” system [47] if thermal data are referenced, and the Hernqvist [13] system if X-ray data are referenced. When appropriate, only the Wille and Lutton system is used.

2. Polymorphic Formation and Transformation in Pure Cocoa Butter

A good description of the procedures necessary to form the six cocoa butter polymorphs is available [50]. The microscopic analyses used in this work also provide insight into the crystal habit of the forms. Each crystal formation procedure required a different method of tempering. Other than rapid cooling of the melt to 0°C or lower, no reference was made by any of the authors regarding the rate of cooling. Neglect of this crystallization parameter or the assumption that the crystallization temperature was most important is one explanation for the lack of reference to cooling rate. In fact, van Malssen et al. [52] found that crystallization temperature, not rate, determined which form would crystallize from the melt. However, their data indicate a possible discrepancy. They stated at 0.25°C/min cooling, β' crystals formed between 22°C and 26°C. At a cooling rate of 1°C/min, the α form crystallized at less than 23°C. Further, at a cooling rate of 6°C/s, both α and γ (sub-α) forms crystallized at less than 3°C. These authors’ observation of no β crystallization after 10 days at 28°C is supported by recent real-time X-ray data, which showed no direct crystallization of β (form VI) from cocoa butter melts [53]. As discussed above, cocoa butter transforms monotonically, from the least stable to the more stable polymorphs. Figure 8 depicts the polymorphic development in cocoa butter.

One unique consequence of the polymorphism of cocoa butter is called fat “bloom.” Investigations of bloom formation were published as early as 1937 [54]. Bloom occurs in chocolate, where it appears as a thin coating or scattered white patches on the surface. Neville et al. [55] attempted to describe the mechanism of bloom formation in terms of physical expansion of melting fat pushing higher melting solid fat to the surface. Full [56] presented a good review of the current understanding of bloom. Three possible mechanisms of bloom formation have been described [57] and are listed in Table 4. Schlichter-Aronhime and Garti [58] argued that because bloom results from a form IV to V conversion and from a V to VI forms and then

### Table 3  Nomenclature and Melting Point (°C) of Cocoa Butter Polymorphs

<table>
<thead>
<tr>
<th>Wille and Lutton [47] Form</th>
<th>mp</th>
<th>Lovegren et al. [49] Form</th>
<th>mp</th>
<th>Hicklin et al. [50] Form</th>
<th>mp</th>
<th>Davis and Dimick [78] Form</th>
<th>mp</th>
<th>Hernqvist [51]* Form</th>
<th>mp</th>
<th>Larson [27]</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>17.3</td>
<td>VI</td>
<td>13.0</td>
<td>I</td>
<td>17.9</td>
<td>I</td>
<td>13.1</td>
<td>sub-α</td>
<td>β'</td>
<td>α or γ</td>
</tr>
<tr>
<td>II</td>
<td>23.3</td>
<td>V</td>
<td>20.0</td>
<td>II</td>
<td>24.4</td>
<td>II</td>
<td>17.1</td>
<td>α</td>
<td>α'</td>
<td>α</td>
</tr>
<tr>
<td>III</td>
<td>25.5</td>
<td>IV</td>
<td>23.0</td>
<td>III</td>
<td>27.7</td>
<td>III</td>
<td>22.4</td>
<td>β'</td>
<td>α + β</td>
<td>β</td>
</tr>
<tr>
<td>IV</td>
<td>27.3</td>
<td>III</td>
<td>25.0</td>
<td>IV</td>
<td>28.4</td>
<td>IV</td>
<td>26.4</td>
<td>β'</td>
<td>β'</td>
<td>β'</td>
</tr>
<tr>
<td>V</td>
<td>33.8</td>
<td>II</td>
<td>30.0</td>
<td>V</td>
<td>33.0</td>
<td>V</td>
<td>30.7</td>
<td>β</td>
<td>β</td>
<td>β</td>
</tr>
<tr>
<td>VI</td>
<td>36.3</td>
<td>I</td>
<td>33.5</td>
<td>VI</td>
<td>34.6</td>
<td>VI</td>
<td>33.8</td>
<td>β</td>
<td>β</td>
<td>β</td>
</tr>
</tbody>
</table>

*From StOSt X-ray diffraction data and PSTP X-ray diffraction data.
Table 4  Possible Mechanisms of Chocolate Fat Bloom Formation

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Blood mechanism</th>
</tr>
</thead>
<tbody>
<tr>
<td>Poor tempering with good cooling and storage</td>
<td>Type IV crystals rapidly transform to type V, resulting in excessive bloom</td>
</tr>
<tr>
<td>Good tempering and good cooling and storage</td>
<td>Type V crystals form and transform to type VI crystals, resulting in variable levels</td>
</tr>
<tr>
<td>Mixed triacylglycerol systems leading to</td>
<td>Liquefied fat recrystallizing on chocolate surface during and after cooling, resulting</td>
</tr>
<tr>
<td>solvation and recrystallization after tempering</td>
<td>in a variable rate of bloom formation</td>
</tr>
</tbody>
</table>

Source: Ref. 57.

conversion, the condition is caused by liquefaction of fat and does not depend strictly on the appearance of a specific polymorphic form. Regardless of the actual mechanism or mechanisms of bloom formation, the polymorphism and phase behavior of cocoa butter underlie the phenomenon.

3. Natural Fat Mixtures and Polymorphism

Although fat mixtures are not used in the confectionery industry alone, this chapter emphasizes the confectionery fat blends specifically used in chocolate and chocolate-like products. Cocoa butter is the primary fat used in chocolate. Its expense has led to the development of other fats, used alone or in combination, to replace some or all cocoa butter in cocoa containing confections. The general term applied to these fats is “confectionery fat.” Two general subclassifications of cocoa butter replacers (CBRs) exist: cocoa butter equivalents (CBE) and cocoa butter substitutes (CBS) [59]. Essentially, a CBE is a mixed fat that provides a fatty acid and triacylglycerol composition similar to those of cocoa butter. A CBS is a fat that provides some of the desired physical characteristics to a confection independent of its dissimilar chemical composition to that of cocoa butter. The development and use of these fats successfully resulted only after the phase behavior of cocoa butter alone and in fat mixtures had been assessed.

CBE fat blends must be tempered, since they will exhibit polymorphism similar to that of cocoa butter. The most stable form is $\beta$. Production of these fats is achieved by blending fractionated and natural fats to achieve a sn-2-oleoyl-disaturated triacylglycerol composition equal to cocoa butter. Ideally, blends of CBE and cocoa butter will be compatible in all proportions. (“Compatibility” refers to the phase behavior of the blend; complete compatibility implies no eutectic effect at any composition.) A thermal eutectic is apparent when the melting point of a blend is lower than that of any of the pure fats in the system.

Unlike CBE, CBS blends have been developed to mimic the hardness and melting properties of cocoa butter only; they are chemically dissimilar to cocoa butter. Palm kernel oil contains a fraction with a triacylglycerol composition that forms a stable $\beta'$-2 polymorph but exhibits physical properties like that of cocoa butter. However, the $\beta$-3 stable form of cocoa butter precludes substantial mixing of cocoa butter and palm kernel oil, since the two fats develop incompatible polymorphic arrangements that lead to softening [60].

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Other mixed-fat systems have been studied in detail. Cocoa butter was found to exhibit only minimal changes in polymorphism when mixed with up to 30% milk fat [41]. Although shifts in the X-ray spectra of these blends were minimal, physical softening of cocoa butter by milk fat and milk fat fractions has been shown [56]. Reddy et al. [61] used DSC to evaluate the polymorphic development and transitions in cocoa butter–milk fat systems. The HMF delayed polymorph transitions, while the LMF facilitated transformation (since the LMF is liquid at normal ambient temperatures). Similar effects in systems of olive oil and cocoa butter have also been observed [62]. It is probable that the liquid fat increases the mobility of the crystal matrix, allowing more rapid conformational changes. The same mechanism may act in LMF cocoa butter blends. The transformation inhibition by HMF supports earlier data showing that hydrogenated milk fat inhibited chocolate fat bloom [63].

4. Effects of Emulsifiers and Other Additives

Schlichter-Aronhime and Garti [58] compared the addition of low percentages of surfactants to cocoa butter to samples of cocoa butter blended with other fats. They noted that fats can change the melting ranges and the number and type of polymorphs, dependent on system compatibility, whereas surfactants may affect the rate of transformation without sensibly altering the crystal lattice. These authors’ generalization of work regarding surfactants and polymorphism stated that surfactants stabilize metastable polymorphs, thereby delaying transformation to the most stable form.

Chocolate bloom was found to be inhibited by sorbitan monostearate [64]. Also, sorbitan tristearate was shown to inhibit the transformation of form V cocoa butter to form VI [58]. However, in the same series of experiments, these authors found that sorbitan tristearate hastened the transformation of the metastable polymorphs to form V. They reported that this occurs as a result of the liquefaction effect of the emulsifier and the liquid-mediated transformation process to form V. They reasoned that form V is stabilized because it transforms to form VI via the solid state, simply by expulsion of trapped liquid. Therefore, the emulsifier promotes only liquid-mediated transformations.

The effectiveness of the emulsifier relative to its state appears conflicting. Schlichter-Aronhime and Garti [58] stated that solid emulsifiers efficiently delay transformations, whereas liquid ones have no effect. However, in another publication the same authors [65] stated that “liquid emulsifiers . . . will enhance the α to β transformation due probably to their weak structure compatibility with tristearin which causes a higher mobility of triacylglycerol molecules.”

Hydrogenated canola oil, which is stable in the β form, can be preserved in the β form by adding 3–5% of 1,2-diacylglycerol. However, 1,3-diacylglycerols were not effective [66–68]. The action of emulsifiers on the polymorphism of fats appears to be related to the physical structure, chemical composition, and thermal properties of both the fat and the additive.

Like addition of emulsifiers to fats, addition to fat of seeding materials, including triacylglycerol seeds, has also been investigated with regard to crystallization rates and polymorphic transitions. The possibility of forming a stable β form in cocoa butter with the addition of preformed seed crystals of cocoa butter and triacylglycerols was investigated [69]. These data showed that a suitable seed must have a chemical composition close to that of the predominant cocoa butter fats. This as well
as more recent efforts [70] showed that the final cocoa butter polymorph depends on the crystallization temperature, not the polymorphic form of the added seed.

The effects of several triacylglycerol seeds on the polymorphism and solidification of cocoa butter in dark chocolate have also been investigated [71,72]. Regarding crystallization rate, it was concluded that thermodynamic stability and crystal structure similarity to cocoa butter are most effective. Specifically, 1,3-distearoyl-2-oleoylglyceride (StOSt) produced greater enhancement than 1,3-dibehenoyl-2-oleoylglyceride (BOB), which produced a much greater enhancement than 1,2,3-tristearoylglyceride (StStSt). It seems that a chain length similarity between the cocoa butter and the crystal seed better facilitates crystal growth. Interestingly, the \( \beta \) form of the StOSt produced a greater crystallization rate than a mixture of \( \beta' \) and \( \beta \). Therefore, although crystallization temperature alone determines the final cocoa butter form, the rate of crystallization may be a function of the polymorphic form of the seed.

5. Intrinsic “Seeding” and Cocoa Butter Crystallization

The crystal morphology of cocoa butter during isothermal crystallization has been detailed [73]. A higher concentration of StOSt was found in crystals that developed early versus the original StOSt concentration in the melt. It was hypothesized that subsequent crystallization occurred on these crystal “seeds.” Such fractional crystallization may produce crystal seeds, rich in StOSt, which have been shown to accelerate cocoa butter crystallization [71,72].

Compounds other than triacylglycerols, found in cocoa butter, have also been investigated for their potential to act as crystal seeds. High-melting crystals isolated from crystallizing cocoa butter melts during the very early stages of crystallization were found to contain very high phospholipid and glycolipid concentrations relative to the original cocoa butter [74,75]. Dimick [76a,76b] proposed that these amphiphilic compounds may associate with the small amount of water in cocoa butter and serve as the nuclei for crystallization (Fig. 12). The phospholipid species found in both the crystal seed and the original butter were identified [11,77]. The data indicated that faster crystallizing cocoa butters contained a relatively high percentage of phosphatidylcholine and phosphatidylglycerol, whereas the slower crystallizing samples had more phosphatidylinositol and significantly less phosphatidylcholine.

The effects on cocoa butter crystallization from simple degumming and those due to added phospholipid seed material were subsequently investigated in the same laboratory [79,80]. Degummed Bahian and Côte d’Ivoire cocoa butters both had significantly slower crystallization rates relative to the untreated butters. Addition of pure \( sn-1,2 \)-distearoylphosphatidylcholine completely inhibited the crystallization of the original butters. However, addition of 0.1% of this phospholipid to the degummed butters increased their crystallization rates to match the untreated samples. After tempering, the original Côte d’Ivoire butter containing pure \( sn-1,2 \)-distearoylphosphatidylcholine had greater solids at 30°C (via NMR) than the untreated butter. Pure \( sn-1,2 \)-dioleoylphosphatidylcholine increased the crystallization rate of the Côte d’Ivoire butter but inhibited the crystallization of the Bahian butter. Also, as with \( sn-1,2 \)-distearoylphosphatidylcholine, added \( sn-1,2 \)-dioleoylphosphatidylcholine in the Côte d’Ivoire butter significantly increased the solids at 30°C. These data show that phospholipids can increase the crystallization rate and enhance the development of the more stable polymorphic form of cocoa butter.
Figure 12  Proposed intermediate structure of high-melting lipid seed crystal illustrating polar core with double chain length saturated triacylglycerol surface [76a,b].
Clearly, the compositional complexity of commercial cocoa butter complicates our understanding of its crystallization and polymorphic transitions. However, the phenomena observed with pure phospholipids in cocoa butter may be of general use in the fats and oils industry with regard to the control and manipulation of solidification during processing.

REFERENCES


I. INTRODUCTION

Interesterification, hydrogenation, and fractionation are three processes available to food manufacturers to tailor the physical, and chemical properties of food lipids [1,2]. At present, roughly one-third of all edible fats and oils in the world are hydrogenated, whereas ~10% are either fractionated or interesterified [3]. Each operation is based on different principles to attain its goal. Fractionation is a physical separation process based on the crystallization behavior of triacylglycerols [4,5]. Hydrogenation, on the other hand, is a chemical process leading to the saturation of double bonds present in fatty acids to harden fats for use as margarine and shortening basestocks. Interesterification, also a chemical process, causes a fatty acid redistribution within and among triacylglycerol molecules, which can lead to substantial changes in lipid functionality. This chapter discusses the application of the theory of chemical interesterification to the production of edible fats and oils.

II. LIPID COMPOSITION

The chemical composition of a fat partly dictates its physical and functional properties [6]. The chemical nature of lipids is dependent on fatty acid structure and distribution on the glycerol backbone. Fatty acids vary in chain length and in the number, position, and configuration of double bonds [7]. Triacylglycerols composed
of saturated fatty acids (e.g., myristic, palmitic, stearic) have high melting points and are generally solid at ambient temperature, whereas triacylglycerols consisting of unsaturated (monoene, polylene) fatty acids (e.g., oleic, linoleic, linolenic) are usually liquid at room temperature. Butterfat, for example, contains ~70% saturated fatty acids, whereas many vegetable oils contain almost exclusively unsaturated fatty acids [4,8].

The fatty acid distribution within naturally occurring triacylglycerols is not random [9,10]. The taxonomic patterns of vegetable oils consist of triacylglycerols obeying the 1,3 random-2-random distribution, with saturated fatty acids being located almost exclusively at the 1,3-positions of triacylglycerols [8,11,12]. Conversely, fats from the animal kingdom (tallow, lard etc.) are quite saturated at the sn-2 position [13].

The industrial applicability of a given fat is limited by its nonrandom distribution, which imparts a given set of physical and chemical properties. The objective of modification strategies, such as chemical interesterification, is the creation from natural fats of triacylglycerol species with new and desirable physical, chemical, and functional properties [14].

III. A BRIEF HISTORY

Interesterification reactions have been knowingly performed since the mid-1800s. The first published mention was by Pelouze and Gélis [15]. Duffy [16] performed an alcoholysis reaction between tristearin and ethanol. Later, Friedel and Crafts [17] generated an equilibrium interchange between ethyl benzoate and amyl acetate. Glyceride rearrangement was also reported by Grün [18], Van Loon [19,20], and Barsky [21]. The first publication demonstrating the chemical interesterification of edible lipids was presented by Normann [22]. Chemical interesterification has been industrially viable in the food industry since the 1940s, to improve the spreadability and baking properties of lard [23,24]. In the 1970s, there was renewed interest in this process, particularly as a hydrogenation replacement for the manufacture of zero-trans margarines. Today it plays a key role in the production of low-calorie fat replacers, such as Proctor and Gamble’s Olestra and Nabisco’s Salatrim or Benefat [25,26].

IV. THE FOUR FACES OF INTERESTERIFICATION

Excellent reviews in the area of chemical interesterification include Sreenivasan [1], Rozenaal [11], Kaufmann et al. [27], Going [28], Hustedt [29], and Marangoni and Rousseau [30].

Interesterification can be divided into four classes of reactions: acidolysis, alcoholysis, glycerolysis, and transesterification [28,31,32].

Acidolysis involves the reaction of a fatty acid and a triacylglycerol. Reactions can produce an equilibrium mixture of reactants and products or can be driven to completion by physically removing one of the reaction products. For example, coconut oil and stearic acid can be reacted to partially replace the short chain fatty acids of coconut oil with higher melting stearic acid [28].

Alcoholysis involves the reaction of a triacylglycerol and an alcohol and has several commercial applications, primarily the production of monoacylglycerols and diacylglycerols. Alcoholysis must be avoided in the interesterification of food lipids,
however, since monoacylglycerols and diacylglycerols are undesirable by-products [28]. Glycerolysis is an alcoholysis reaction in which glycerol acts as the alcohol [31].

Transesterification is the most widely used type of interesterification in the food industry. Hence, we concentrate on this reaction. Figure 1 shows the effects of interesterification on the fatty acid distribution of a putative triacylglycerol (1-stearoyl-2-oleoyl-3-linoleoyl glycerol) (SOL). In sequence, the ester bonds linking fatty acids to the glycerol backbone are split; then the newly liberated fatty acids are randomly shuffled within a fatty acid pool and reesterified onto a new position, either on the same glycerol (intraesterification) or onto another glycerol (interesterification) [1]. For reasons involving thermodynamic considerations, intraesterification occurs at a faster rate than interesterification [33]. Once the reaction has reached equilibrium, a complex, random mixture of triacylglycerol species is obtained (Fig. 1).

The extent of the effects of interesterification on the properties of a fat will depend on the fatty acid and triacylglycerol variety of the starting material. If a single starting material (e.g., palm stearin) is randomized, the effects will not be as great as if a hardstock is randomized with a vegetable oil [34]. Furthermore, if a material has a quasi-random distribution prior to randomization (e.g., tallow), randomization will not lead to notable modifications.

The interesterification reactions consists of three main steps: catalyst activation, ester bond cleavage, and fatty acid interchange. We now examine each subject in detail.

V. INTERESTERIFICATION CATALYSTS

A. Is a Catalyst Necessary for Interesterification?

Interesterification can proceed without catalyst at high temperatures (~300°C); the desired results are not obtained, however, because equilibrium is slowly attained at such temperatures, and isomerization, polymerization, and decomposition reactions can occur [11,28,35]. In fact, polymerization has been shown to occur at 150°C [36]. Addition of catalyst significantly lowers reaction temperature and duration [37]. Other important considerations include the type and concentration of catalyst [38].

B. Available Catalysts for Interesterification

There are three groups of catalysts (acids, bases, and their corresponding salts and metals), which can be subdivided into high and low temperature groups [27]. High temperature catalysts include metals salts such chlorides, carbonates, oxides, nitrates, and acetates of zinc, lead, iron, tin, and cobalt [39]. Others include alkali metal hydroxides of sodium and lithium [40]. Most commonly used are low temperature catalysts such as alkylates (methylate and ethylate) of sodium and sodium/potassium alloys; however, other bases, acids, and metals are also available [41]. Alkylates of sodium are simple to use and inexpensive, and only small quantities are required. Furthermore, they are active at low temperatures (<50°C). This last characteristic allows their use for directed interesterification [1,42].

C. Precautions

Performing a chemical interesterification reaction is a relatively straightforward process. However, a 100% reaction yield is never attainable [43]. Volatile fatty acid
alkyl esters, formed in stoichiometric yields with the catalyst during the reaction, must be washed out, and a small amount of partial acylglycerols is always produced [44]. Trace amounts of moisture will inactivate alkylate catalysts by producing the corresponding alcohols. Hence, the fat or oil should contain less than 0.01% (w/w) water [29]. Free fatty acids and peroxides also impair catalyst performance, and levels should be maintained as low as possible, preferably below 0.1% and 1.0% (w/w), respectively [37]. The fat should be well refined, dried, and heated (120–150°C) under a nitrogen blanket before addition of catalyst [14]. Finally, sodium alkylates are toxic, highly reactive materials that should be handled with care. Their shelf life is a few months [11].

With a dry oil devoid of impurities, only trace amounts of catalyst [<0.4% (w/w)] are required [14]. Catalyst concentration should be minimized to prevent excessive losses due to saponification [42]. Experience has shown that above 0.4% catalyst, the addition of each additional 0.1% of catalyst results in the loss of ~1% neutral fat [29]. Konishi et al. [45], however, observed that ester interchange between soybean oil and methyl stearate in hexane was improved by using 10% (w/w) sodium methoxide in hexane. Proportions between 0.1% and 4.0% led to similar amounts of ester interchange.

It is also necessary to use the catalyst in a form that is easily and completely dispersed [46]. For example, if Na/K catalysts are not finely dispersed in a suitable solvent, a violent reaction with residual moisture may occur at the catalyst surface, followed by splitting of surrounding fat molecules to form a coating of soap. The heat generated by such a reaction is enough to decompose triacylglycerols and cause local charring [28].

Figure 1: Triacylglycerol formation during interesterification: S, stearoyl; O, oleoyl; L, linoleoyl. (Adapted from Ref. 1.)
D. The “Real” Catalyst

The real catalyst is believed to be a metal derivative of a diacylglycerol, and the aforementioned catalysts are most likely its precursor [11, 42]. Upon catalyst addition to the lipid, a reddish brown color slowly develops (within a few minutes, depending on the application and reaction conditions) in the mixture, indicating the activation of the presumed true catalyst.

Some workers time the interesterification reaction from the appearance of the reddish brown color; others simply time the reaction from the moment of catalyst addition. Because it is impossible to predict reaction onset, it is difficult to obtain a partial interesterification. Most reactions are conducted until an equilibrium has been reached. Reaction times are longer in industrial settings, because the catalyst must be totally homogenized within the fat [43]. Preactivation is unnecessary if the catalyst is predissolved prior to addition to the substrate [35]. Placek and Holman [47] incorrectly attributed the induction period to the interaction between the catalyst and impurities. While impurities are sometimes present, the induction period is not strictly due to their presence; rather, it is due to catalyst activation. As stated by Coenen [41] and many others, the activation energy for the catalyst is higher than for the reaction. A preactivation of 15 minutes has been found to accelerate the reaction itself [45]. Interestingly, Hustedt [29] stated that once the brown intermediate had appeared in the reaction mixture, interesterification was complete. No basis was given for this statement.

E. Reaction Termination

The interesterification reaction is allowed to continue for a predetermined time period and is stopped with addition of water and/or dilute acid. Going [28] described three patents dealing with catalyst removal techniques for minimizing fat loss. Generally, most of the catalyst can be washed out with water to a separate salt, or a soap-rich aqueous phase. Alternatively, reaction with phosphoric acid results in a solid phosphate salt, which can be filtered out. Both these methods result in substantial fat loss. A technique has been developed that minimizes loss by addition of CO$_2$ along with water. The system becomes buffered with sodium carbonate at a pH low enough to not split the fat [48].

VI. REACTION MECHANISMS

The exchange of fatty acids between triacylglycerol hydroxyl sites does not occur directly but via a series of alcoholysis reactions involving partial acylglycerols [49]. The proposed mechanisms of chemical interesterification depend on the inherent properties of the triacylglycerol ester carbonyl group (C=O). The carbonyl carbon is particularly susceptible to nucleophilic attack due to electronic and steric considerations. The electronegative oxygen pulls electrons away from the carbonyl carbon, leading to a partial positive charge on the carbon, and also increases the acidity of hydrogens attached to the carbon at a position $\alpha$ to the carbonyl group (Fig. 2).

Steric considerations also come into play. The carbonyl carbon is joined to three other groups by $\sigma$ bonds ($sp^{2}$ orbitals); hence they lie in a flat plane, 120° apart. The remaining $p$ orbital from the carbon overlaps with a $p$ orbital from the oxygen, forming a $\pi$ bond. This flat plane and the absence of neighboring bulky
groups permits easy access for nucleophiles to approach and react with the carbonyl carbon.

The transition state of the reaction is a relatively stable tetrahedral intermediate with a partial negative charge on the oxygen. As the reaction progresses, a group leaves and the structure reverts to the planar carbonyl structure. Strong evidence supports the cleavage of the carbonyl carbon–oxygen bond as the mechanism for the release of the leaving group.

For acid-catalyzed nucleophilic acyl substitution, a hydrogen easily associates with the carbonyl oxygen owing to the polarized nature of the carbonyl function and the presence of free electron pairs on the oxygen, imparting a positive charge to this atom [50]. The carbonyl carbon is then even more susceptible to nucleophilic attack, since oxygen can accept \( \pi \) electrons without gaining a negative charge. Acid-catalyzed interesterification is not discussed further because it is not used for the chemical interesterification of food lipids.

A. Carbonyl Addition Mechanism

In alkaline conditions encountered during interesterification, the catalyst (which is nucleophilic) attacks the slightly positive carbonyl carbon at one of three fatty acid–glycerol ester bonds and forms a tetrahedral intermediate. The fatty acid methyl ester is then released, leaving behind a glyceryl anion (Fig. 3A). Kinetics of base-catalyzed hydrolysis of esters shows that the reaction is dependent on both ester and base concentration (second-order kinetics). This newly formed glyceryl anion is the nucleophile for subsequent intra- and intermolecular carbonyl carbon attacks, which continue until a thermodynamic equilibrium has been reached (Fig. 4).
During an attack, a new triacylglycerol is not necessarily formed. The transition complex (glycerylate + fatty acid) will decompose, either to regenerate the original species and active catalyst or to form a new triacylglycerol and a new active catalyst ion. This process continues until all available fatty acids have exchanged positions and an equilibrium composition of acylglycerol mixture has been achieved [1]. Support for this mechanism was provided by Coenen [41] who presented the kinetics between a simple mixture of S3 and U3. The kinetics were described with six possible reactions between various triacylglycerol and diacylglycerol anions, with a rate con-
stant 3k (Fig. 5). Not all possible exchanges produced a net change in triacylglycerol composition, leading to 2k and k rate constants.

**B. Claisen Condensation**

In Claisen condensations, the sodium methoxide removes an acidic hydrogen from the carbon α to the carbonyl carbon, yielding an enolate ester [23]. This reaction produces a carbanion, a powerful nucleophile (Fig. 3B). This nucleophile will attack carbonyl groups, forming a β-keto ester intermediate and a glycerylate. The glycer-
ylate is now free to attack other carbonyl carbons and exchange esters intra- and 
intermolecurally (Fig. 6). Once this carbanion has been created, the same consider-
ations as for the usual carbonyl carbon chemistry apply.

VII. RANDOM AND DIRECTED INTERESTERIFICATION

A. Random Interesterification

Interesterification reactions performed at temperatures above the melting point of the 
highest melting component in a mixture result in complete randomization of fatty 
acids among all triacylglycerols according to the laws of probability [28,51].

The energy differences between the various combinations of triacylglycerols 
are insignificant and do not appear to lead to fatty acid selectivity [52]. Hence, 
random interesterification is entropically driven (randomization of fatty acids among 
all possible triacylglycerol positions) until an equilibrium is reached [41].

In ester–ester interchange, the fatty acid distribution is theoretically fully ran-
domized, meaning that the resulting triacylglycerol structure can be predicted from 
the overall fatty composition of the mixture (Table 1) [11].

In Table 1, \( a \), \( b \), and \( c \) are the molar concentrations of fatty acids \( A \), \( B \), and \( C \). 
\( AAA \), \( AAB \), and \( ABC \) are triacylglycerols composed of one, two, or three different 
fatty acids, respectively. For \( AAB \), there are three possible isomers, whereas for \( ABC \) 
there are six. For example, 1-stearoyl-2-oleoyl-3-linoleoyl glycerol results in the 
following fully randomized equilibrium mixture:

\[
\begin{align*}
S_3 + U_2 ONa & \rightleftharpoons \frac{3k}{k} SU_2 + S_2 ONa \\
U_3 + S_2 ONa & \rightleftharpoons \frac{3k}{2k} S_2 U + U_2 ONa \\
SU_2 + U_2 ONa & \rightleftharpoons \frac{3k}{k} U_3 + SUONa \\
S_2 U + S_2 ONa & \rightleftharpoons \frac{3k}{2k} S_2 + SUONa \\
S_2 U + U_2 ONa & \rightleftharpoons \frac{3k}{k} SU + SUONa \\
SU_2 + S_2 ONa & \rightleftharpoons \frac{3k}{k} S_2 U + SUONa
\end{align*}
\]

Figure 5 Kinetics of interesterification via the carbonyl reaction mechanism (S, SS, SSS 
and U, UU, UUU: mono-, di-, and trisaturated and unsaturated, respectively). (Adapted from 
Ref. 41.)
Figure 6  Reaction mechanism for the chemical inter- and intraesterification of two triacylglycerols via the Claisen condensation mechanism. (Adapted from Ref. 30.)
Table 1  Theoretical Triacylglycerol Compositions After Complete Interestesterification of \( n \) Fatty Acids (A, B, C, D, \ldots) with Molar Fractions \( a, b, c, d, \ldots \)

<table>
<thead>
<tr>
<th>Type</th>
<th>Quantity</th>
<th>Proportion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Monoacid (AAA, BBB, \ldots)</td>
<td>( n )</td>
<td>( a^3, b^3, \ldots )</td>
</tr>
<tr>
<td>Diacid (AAB, AAC, \ldots)</td>
<td>( n(n-1) )</td>
<td>( 3a^2b, 3a^2c, \ldots )</td>
</tr>
<tr>
<td>Triacid (ABC, DEF, \ldots)</td>
<td>( \frac{n(n-1)(n-2)}{6} )</td>
<td>( 6abc, 6def, \ldots )</td>
</tr>
<tr>
<td>Total</td>
<td>( \frac{n(n+1)(n+2)}{6} )</td>
<td></td>
</tr>
</tbody>
</table>

Source: Adapted from Ref. 11.

SSS, 3.7%  SOO, 11.1%
OOO, 3.7%  SLL, 11.1%
LLL, 3.7%  OOL, 11.1%
SSO, 11.1% OLL, 11.1%
SSL, 11.1% SOL, 22.2%

Gavriilidou and Boskou [53] found that a random distribution was obtained after chemical interesterification of olive oil–tristearin blends. They observed that trisaturate and triunsaturate proportions decreased markedly, whereas proportions of SSU and SUU increased (Table 2).

Not all workers agree that chemical interesterification is a purely random process. Kuksis et al. [54] found that the triacylglycerol composition of rearranged butter and coconut oils approached random distribution but deviated from true random distribution, even when experimental error was accounted for. This result was attributed to differences in the reactivity of the fatty acids and to possibly different esterification rates of the inner and outer hydroxyl sites on the glycerol backbone.

B. Batch Interestesterification

Random interesterification can be accomplished in either batch or continuous mode. A typical batch reactor (Fig. 7) consists of a reaction vessel fitted with an agitator, heating/cooling coils, nitrogen sparger, and vacuum pump [3,14,28]. In a batch process, the raw lipid is heated to 120–150°C under vacuum in the reaction vessel to remove any trace of moisture [1,29]. As mentioned, moisture and peroxides deactivate the catalyst. Following the drying step, the mixture is cooled to 70–100°C. Catalyst is sucked into the reaction vessel and disperses to form a white slurry. The reaction is allowed to proceed for 30–60 minutes. When completion has been confirmed by analysis, the catalyst is neutralized in the reaction vessel. Processing losses can be minimized by using as little catalyst as possible and neutralizing with phosphoric acid or CO\(_2\) prior to addition of water.

C. Continuous Interestesterification

During continuous random interesterification, the fat is flash-dried and catalyst is continuously added. The fat then passes through elongated reactor coils with res-
Table 2  Triacylglycerol Makeup for Olive Oil–Tristearin Blends Before and After Interesterification

<table>
<thead>
<tr>
<th>Species&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Olive oil–tristearin blend</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>75–25% (w/w)</td>
</tr>
<tr>
<td>SSS</td>
<td>Initial 25.1</td>
</tr>
<tr>
<td></td>
<td>Randomized 4.4</td>
</tr>
<tr>
<td>SSU</td>
<td>Initial 3.4</td>
</tr>
<tr>
<td></td>
<td>Randomized 24.1</td>
</tr>
<tr>
<td>UUS</td>
<td>Initial 23.4</td>
</tr>
<tr>
<td></td>
<td>Randomized 44.3</td>
</tr>
<tr>
<td>UUU</td>
<td>Initial 46.9</td>
</tr>
<tr>
<td></td>
<td>Randomized 27.2</td>
</tr>
</tbody>
</table>

<sup>a</sup>S, saturated; U, unsaturated.

Source: Ref. 53.

idence time determined by the coil length and the flow rate of the oil. The catalyst is then neutralized with water, separated from the oil by centrifugation, and dried [14].

Rozenaal [11] mentioned a continuous interesterification process in which a solution of sodium hydroxide and glycerol in water was used as precatalyst. Heated oil was mixed with the catalyst solution and subsequently spray-dried in a vacuum drier to obtain a fine dispersion and to remove the water. The reaction could be carried out in a coil reactor at 130°C. With this setup, the reaction took only a few minutes.

D. Regioselectivity in Interesterification

Elegant work by Konishi et al. [45] demonstrated that chemical interesterification can be regioselective. Sodium methoxide–catalyzed ester interchange between soybean oil and methyl stearate in hexane, at 30°C, revealed that fatty acid interchange at sn-1,3 positions progressed 1.7 times faster than at the sn-2 position after 24 hours of reaction.

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Cast et al. [55] demonstrated the regioselectivity of chemical interesterification in the presence of phase transfer catalysts (tetraalkyl ammonium bromides), with trilaurin (LaLaLa) and methyl palmitate in the presence of sodium methoxide. Under normal conditions, interesterification resulted in 17.7% LaLaP and 82.3% unreacted LaLaLa. Using tetrahexyl ammonium bromide, 11.9% LaLaP, 6.2% LaPP, and 81.2% LaLaLa were obtained. Seven quaternary ammonium salts were tested for their effect on the reaction: tetraethyl, tetrapropyl, tetrabutyl, tetrapentyl, tetrahexyl, and tetraheptyl ammonium bromides. The amount of LaLaOH diacylglycerol increased as the chain length increased up to tetrahexyl ammonium bromide (Fig. 8). The amounts of LaPOH and PPOH diacylglycerol species increased up to tetraheptyl ammonium bromide. Reactions were performed for 48 hours, with a fivefold increase in the amount of LaPOH diacylglycerols compared to the amount after 1 hour of reaction.

Most importantly, following lipase hydrolysis and subsequent 2-monoacylglycerol isolation, it was discovered that the reaction between trilaurin and methyl palmitate in the presence of tetrapentyl ammonium bromide contained 49% lauric acid and 51% palmitic acid, which represents an enrichment factor for the 2-position of 1.51 times. Hence, under these conditions, chemical interesterification was not a random process.

**E. Directed Interesterification**

If the interesterification reaction is carried out at temperatures below the melting point of the highest melting component (most likely a trisaturated triacylglycerol species), the end result will be a mixture enriched in this component. This was first reported by Eckey [35] who, for the interesterification of lard, discovered that certain

**Figure 8** Composition of diacylglycerol content versus phase transfer catalyst chain length for sodium methoxide–catalyzed interesterification of trilaurin and methyl palmitate in the presence of seven quaternary ammonium salts. PPOH (○); LaPOH (●); LaLaOH (□). (Adapted from Ref. 55.)
catalysts were active below the melting point of the fat and that the reaction reached equilibrium within 30 minutes.

During directed interesterification, two reactions take place simultaneously. As the trisaturate is produced by interesterification, it crystallizes and falls from solution. Then, to regain equilibrium, the reaction equilibrium in the remaining liquid phase is pushed toward increased production of the crystallizing trisaturate [42,47,56]. Crystallization continues until all triacylglycerols capable of crystallizing have been eliminated from the reaction phase [43].

Early developments in the area of directed interesterification showed that the following factors determine the effectiveness of the reaction [47,56]:

- Interesterification rate in the liquid phase
- Rate of heat removal
- Fat crystal nucleation rate
- Trisaturate crystallization rate out of liquid phase

The rate of interesterification is an important factor, as the trisaturates will precipitate out of solution only as quickly as they are formed.

Fat crystallization generates heat. Removal of this heat is hindered by the poor conductivity of fat and the low convection in viscous or plastic media. Heat removal directly affects the nucleation rate. Rapid cooling to temperatures much below the melting point of trisaturate increases the nucleation rate, hence crystallization. Trisaturate crystallization is also hindered by the viscosity of the lipid phase. Gently yet thorough agitation is helpful in speeding up crystallization.

For directed interesterification, Na/K alloy is the catalyst of choice, given its low-temperature activity compared to that of the metal alone or that of the alkylates [28,47]. Typically, the alloy is continuously metered in by a pump and well dispersed by means of a high shear agitator to provide the proper catalyst particle size, ensuring optimal activity. Initially, the fat is at least partially randomized at temperatures above the melting point of the highest melting triacylglycerol. When the fractional crystallization approach is used, the fat/catalyst slurry is chilled in conventional scraped-wall heat exchangers to specific temperatures in a series of steps designed to maintain the directed fractional crystallization process. Once chilled, the mixture is held for a period of time under gently agitation to achieve the desired degree of crystal formation. Enhancements of the procedure include stepwise reduction of temperature and the use of temperature cycling [57,58].

Directed interesterification can be used to increase the solid fat content without affecting unsaturated fatty acids. Periodic drops in temperature accelerate the reaction. Kattenberg [58] applied this knowledge to interesterification of sunflower oil and lard blends and accelerated the reaction by a factor of 3. In another study, various oils, after directed interesterification, were chilled at 15°C for various durations (30–180 min), then subjected to further reaction at 23°C for 12–168 hours. These treatments influenced solid fat content of the final product [59].

The effects of directed interesterification on cottonseed oil were reported by Eckey [35]. Cottonseed oil contains 25% saturated fatty acids. With random interesterification, only 1.5% trisaturates was obtained, whereas directed interesterification led to the production of 19% trisaturates.

A review by Huyghebaert et al. [60] showed that the directed interesterification of an SOL mixture resulted in the following proportions:
Solid: SSS, 33.3%
Liquid: OOO, 8.3%
OOL, 24.99%
OLL, 24.99%
LLL, 8.3%

The segregation of saturated fatty acids into trisaturated species is necessarily accompanied by a corresponding tendency for unsaturated fatty acids to form triunsaturated species [47].

VIII. KINETICS OF CHEMICAL INTERESTERIFICATION

Random interesterification is usually conducted until equilibrium has been reached. There are many conflicting reports in the literature concerning interesterification reaction rates. Coenen [41] stated that once a sufficient concentration of catalyst in solution had been reached in the reaction mixture, the actual interesterification reaction was extremely fast, requiring only a few minutes, unless operations had to proceed at very low temperatures. The kinetics were modeled in several ways to support this theory. The first example was a model system consisting of short chain fatty esters (C8, C10) of ethylene glycol (Fig. 9). The induction period was long, yet the reaction itself was rapid, even at 32°C. In the second example, interesterification of palm oil was evaluated using solid fat content determinations (Fig. 10). The reaction rate was faster at higher temperatures. These data confirm that an activation period is indeed required and agree with Weiss et al. [24] and Rozenaal [11], who reported that the catalyst formation phase was longer than the interesterification

Figure 9  Theoretical interesterification kinetics of glycol esters of C8 and C10 as a function of time and temperature. 42°C (○); 37°C (●); 32°C (□). (Adapted from Ref. 41.)
reaction, since the activation energy was higher for catalyst formation than for the interesterification reaction itself.

Lo and Handel [61] observed that interesterification of soybean oil and beef tallow was complete after 30 minutes (Fig. 11). Reaction completion was determined by lipase hydrolysis analysis. Results by Konishi et al. [45] showed that in certain cases the interesterification reaction can progress for as long as 24 hours, even with catalyst preactivation. Thus, depending on conditions, randomization can proceed for many hours.

Other factors that may influence interesterification onset include agitation intensity, catalyst particle size, and temperature. Studies by many, including Konishi et al. [45], Laning [14], and Wiedermann et al. [23] have shown that interesterification kinetics are temperature dependent (Fig. 12).

IX. ASSESSING THE EFFECTS OF INTERESTERIFICATION ON LIPID PROPERTIES

Fats and oils are usually modified to attain a certain functionality, such as improved spreadability, a specific melting point, or a particular solid fat content–temperature profile. However, changes in triacylglycerol structure may constitute the purpose of the reaction, as in the synthesis of a particular structure. For that purpose, the fatty acid distribution constitutes the reaction goal. Methods described to assess physical properties include cloud point, Mettler dropping point, pulsed nuclear magnetic resonance, differential scanning calorimetry, cone penetrometry, X-ray diffraction, and polarized light microscopy. Chromatographic methods include thin-layer chromatography (TLC), high pressure liquid chromatography (HPLC), and gas–liquid chro-
Figure 11  Changes in the fatty acid distribution at the sn-2 position during random interesterification of a 60:40 (% w/w) soybean oil–beef tallow mixture: ○, 16:0; ●, 18:0; □, 18:1; ■, 18:2; △, 18:3. (Adapted from Ref. 61.)

Figure 12  Influence of temperature on the interesterification reaction rate with glycerol/NaOH catalyst. (Adapted from Ref. 14.)
matography (GLC). Other methods not discussed include mass spectroscopy [62] and stereospecific lipase hydrolysis [63,64].

### A. Physical Properties

Physical properties can be determined by examining thermal characteristics, rheological characteristics, or crystal habit.

#### 1. Cloud Point

The cloud point, or temperature at which crystallization is induced, producing a crystal cloud, is one of the older indices used to study physical properties of fats. Eckey [35] monitored the change in cloud point during cottonseed oil interesterification. Generally, randomization increased the cloud point quickly at first and then more slowly until an increase of 13–15°C was reached, after which no change was observed, regardless of reaction duration. For Placek and Holman [47], a study of cloud point indicated the extent of interesterification of lard.

#### 2. Dropping Point

The Mettler dropping point is a simple yet effective method of measuring the effect of interesterification on fats. In this procedure, liquefied fats are crystalized in sample cups and subsequently heated until they begin flowing under their own weight. Kaufmann and Grothues [65] performed a thorough study of the dropping points of hardstock and vegetable oil mixtures as an indicator of catalyst activity. Laning [14] demonstrated the effect of chemical interesterification on palm, palm kernel, and coconut oils (Table 3). A reduction in dropping point for saturated palm kernel oil (PKO) and saturated coconut oil was due to the lower average molecular weight of the triacylglycerol in the randomized fat. The reduction in dropping point reported for randomized PKO was due to an increase of triacylglycerol species with intermediate degrees of unsaturation. Cho et al. [66] used the dropping point as an indicator of the measure of reaction equilibrium. A blend of 70% hydrogenated canola oil, 10% palm stearin, and 20% canola oil had an initial dropping point of 37°C, which dropped to 35°C following 5 minutes of reaction and to 32°C after 20 minutes, remaining constant thereafter. List et al. [67] used dropping point as a verification of interesterification completion in the preparation of “zero-trans” soybean oil margarine base stock. Rousseau et al. [34] examined the effect of chemical interesteri-

<table>
<thead>
<tr>
<th>Oil</th>
<th>Before treatment</th>
<th>Random treatment</th>
<th>Directed treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Palm oil</td>
<td>39.4</td>
<td>42.7</td>
<td>51.1</td>
</tr>
<tr>
<td>Palm kernel (PKO)</td>
<td>28.3</td>
<td>26.9</td>
<td>30.0</td>
</tr>
<tr>
<td>Coconut oil (CO)</td>
<td>25.5</td>
<td>28.2</td>
<td>—</td>
</tr>
<tr>
<td>Saturated PKO</td>
<td>45.0</td>
<td>34.4</td>
<td>—</td>
</tr>
<tr>
<td>Saturated CO</td>
<td>37.8</td>
<td>31.6</td>
<td>—</td>
</tr>
</tbody>
</table>

*Source: Ref. 14.*
fication and blending on butterfat–canola oil and found that a linear increase in the proportion of canola oil did not lead to a linear reduction in dropping point.

3. Nuclear Magnetic Resonance

The amount of solid triacylglycerols in a lipid sample can be determined by means of NMR techniques. Laning [14] demonstrated the effect of chemical interesterification on palm, palm kernel, and coconut oils. Random interesterification resulted in modest solid fat content (SFC) changes, while directed interesterification produced more significant increases, attributable to the increase in trisaturated triacylglycerols. Generally speaking, interesterification results in more linear profiles, owing to the greater variety of triacylglycerol species [68].

Blending of butterfat with canola oil produced slight changes in the solid fat content of butterfat–canola oil blends, as exemplified by a contour profile (Fig. 13) [34]. No changes greater than ±6% were evident. The biggest increase in solid fat content produced by interesterification was of butterfat at 25°C; SFC “valleys” were present for the 80% butterfat–20% canola oil blend at 15°C, while the largest decreases were present for the 40% butterfat–60% canola oil blend at 10°C.

4. Differential Scanning Calorimetry

DSC is used to measure the melting or crystallization profile and accompanying changes in enthalpy of fats. Rost [69] described the directed interesterification of palm oil. Calorimetry results indicated that the melting thermogram for noninteresterified palm oil consisted of two main peaks centered around 10°C and 19°C. Di-
rected interesterification of palm oil led to a broader melting profile with no distinct peaks. Rossell [70] studied the effects of chemical interesterification on palm kernel oil crystallization. Randomization did not alter the shape of the crystallization curve; only peak temperatures were slightly lower. Because of the wide range of triacylglycerols that must be packed into fat crystals, interesterified fats generally show simpler melting curves with less polymorphism upon chemical interesterification [68].

Zeitoun et al. [71], who examined interesterified blends of hydrogenated soybean oil and various vegetable oils (1:1 w/w ratio), found that each oil influenced the melting and crystallization behavior of the interesterified blends differently as a result of initial variations in oil composition. Rousseau et al. [34] found that chemical interesterification of butterfat–canola oil blends also led to simpler, more continuous melting profiles. However, overall changes were minimal.

5. Cone Penetrometry

This is a rapid yet empirical method used in the evaluation of fat texture and rheology [72]. Jakubowski [43] found that interesterification doubled penetration depth of a blend of 35–65% tallow–sunflower oil blends, at 15°C. Rousseau et al. [73] reported that interesterification substantially decreased the hardness index of blends of butterfat and canola oil. Other rheological measurements include viscoelasticity measurements [73].

6. X-Ray Diffraction

The polymorphic behavior of fats is important in many food systems (fat spreads, chocolate, etc.) [52]. Fat spread crystals exist as one of three primary forms: α, β', and β. The β modification is to be avoided in fat spreads because it results in a sandy texture [72]. The β' crystals are the most desirable form. Chemical interesterification alters the crystal morphology and structure of fats. Larsson [74] stated that a greater variety of fatty acids hinders β-crystal formation. Hence, upon interesterification of butterfat, which normally consists of a predominance of β'-crystals and a slight proportion of β-crystals, the latter disappeared upon triacylglycerol randomization [68, 75, 76].

List et al. [77], while working with margarine oils, found that chemical interesterification and blending of vegetable oils and hydrogenated hardstock of soybean oil or cottonseed oil resulted in β'-crystal polymorphs.

Hernqvist et al. [52] interesterified mixtures of tristearin, triolein, and trielaidin. These mixtures were chosen to produce model systems for vegetable oil blends used in margarine. Polymorphic transitions of interesterified blends were studied, and depending on the blend, two to four polymorphs (sub-α, α, β', or β) were observed.

7. Polarized Light Microscopy

The morphology of the crystals comprising the three-dimensional fat crystal network is largely responsible for the appearance and texture of a fat and exerts a profound influence on its functional properties. Interesteification leads to noticeable modifications in crystal morphology, which can be examined in great detail with polarized light microscopy [78]. Prior to interesterification, lard consists of large crystals promoting graininess. Following interesterification, tiny delicate crystals, typical of the β' polymorph, are present [79]. Becker [80] performed an in-depth study on the
influence of interesterification on crystal morphology of binary and ternary mixtures of trilaurin, triolein, and tristearin. He also found that fat crystals following interesterification were smaller than before randomization and had different morphologies. A study of butterfat-canola oil blends revealed that gradual addition of canola oil led to gradual spherulitic aggregation of the crystal structure [75].

B. Chemical Properties

Changes in physical properties provide an arbitrary measure of interesterification structural modifications but give no real information on the compositional changes. Following these changes can be difficult unless simple substances are used [33]. Studies on molecular rearrangement of triacylglycerol species provide a true indication of the chemistry of interesterification. The chemistry of interesterification can be followed with different chromatographic techniques: TLC, HPLC, and GLC.

Freeman [33], who examined the changes in monounsaturated triacylglycerols during the course of interesterification with TLC, found that intraesterification occurred at a faster rate than the general randomization that results from interesterification.

Chobanov and Chobanova [57] made extensive use of TLC to study the alteration in composition of 10 triacylglycerol groups during the monophasic interesterification of mixtures of sunflower oil with lard and tallow.

Parviainen et al. [81] studied the effects of randomization on milk fat triacylglycerol; they found an $S_{2}U$ decrease in C36 and C38 species (45% and 52%, respectively) and an increase in trisaturated C44–C50 species. This combination led to a broader crystallization range and higher SFCs at temperatures above 25°C.

Herslöf et al. [82] used reversed phase HPLC and GC to analyze the interesterification reaction between fatty acid methyl esters and trilaurin and found that the theoretical and experimental compositions for the interesterified systems matched.

Rossell [70] measured the evolution in triacylglycerol species following chemical interesterification of palm kernel oil by means of GLC.

Huyghebaert et al. [60] and Rousseau et al. [34] used GLC to follow the evolution of butterfat triacylglycerol species as a result of interesterification. Typical results are shown in Figure 14.

X. APPLYING INTERESTERIFICATION TO FOOD LIPIDS

Chemical interesterification is used industrially to produce fats and oils used in margarines, shortenings, and confectionery fats [60]. Due to legislation and for economic reasons, interesterification is a more common process in Europe than in North America. It is popular for many reasons. For example, little in the way of chemical properties is affected, and the fatty acid distribution is changed but the fatty acids’ inherent properties are not. Moreover, unsaturation levels stay constant and there is no cis-trans isomerization [36,43]. Interesterialization can improve the physical properties of fats and oils. Similar changes in physical properties may be obtained by means of blending, fractionation, or hydrogenation. Production costs, market prices, or raw material and nutritional concerns will determine the process to be used. Applications described include lard, margarines, palm oil and palm kernel oil, milk fat, and fat substitutes.
A. Shortening

Chemical interesterification has been successfully used for decades to improve the physical properties of lard. Ordinary lard has a grainy appearance, a poor creaming capacity, and a limited plastic range, which is not improved by plasticizing in a scraped-surface heat exchanger [43,47]. Addition of a hard stock plus plasticizing helps in these respects, but the product develops an undesirable graininess during storage [83]. Chemical interesterification halves the solid fat content of the lard at 20°C, improves the plastic range of the fat considerably, and prevents the development of graininess, which is due to the large proportion (64%) of palmitic acid at the sn-2 position [83,84]. This improvement in plasticity and stability is due to alterations in the polymorphic behavior, with interesterified lard crystallizing in a β'-2 form, characteristic of hydrogenated vegetable oil shortenings [83].

Chemically, the β'-3-tending disaturated OPS (1-oleoyl-2-palmitoyl-3-stearoylglycerol) (large crystals responsible for lard graininess) are exchanged for a mixture of disaturated triacylglycerols, with a lower melting point and greater intersolubility; the sn-2 palmitic acid concentration drops from 64% to 24% promoting β' behavior [83]. A detectable morphological change that accompanies these chemical changes is an increase in the relative proportion of small fat crystals [85].

Duterte [86] mentioned that the crystalline modifications were observed prior to the theoretical completion of randomization. Production of fine crystals extends lard’s plastic range and gives it a smooth appearance [47]. Random interesterification helps to resolve the graininess problem, yet the limited plastic range problem is not fully resolved. The S,U triacylglycerols in randomized lard give little plasticity at higher temperature. Directed interesterification resolves the plastic range problem [56].
B. Margarines

In the manufacture of margarine, the object is to produce a fat mixture with a steep solid fat content curve to obtain a stiff product in the refrigerator that nevertheless spreads easily upon removal and melts quickly in the mouth. It should crystallize as a $\beta'$ polymorph [87]. Depending on oil costs and availability, different treatments can be used.

As an alternative to hydrogenation for the production of margarine, Lo and Handel [61] chemically interesterified blends of 60% soybean oil with 40% beef tallow. Final results indicated properties similar to those of commercial tub margarine oil. Yet the interesterified blend contained less polyunsaturated fatty acids and more saturated fatty acids than commercial margarine oil.

According to Sonntag [31], short and medium chain fatty acids (C6–C14) have good melting properties whereas long chain fatty acids (C20–C22) can provide stiffening power in margarine. Acids of these two types can be combined with interesterification to produce triacylglycerols that provide blends with good spreadability, high temperature stability, and a pleasant taste.

Margarine oil with high proportions of lauric acid has a low melting point and narrow plastic range, which leads to a margarine that is hard in the fridge but partly melts at room temperature [87]. Decreasing the lauric acid concentration can rectify this problem of extremes. For example, coconut oil can be interesterified with an oil such as palm, and 60% of the interesterified mixture then blended with 40% oil, such as sunflower oil.

In the manufacture of zero-trans margarines, chemical interesterification of soybean oil-soy trisaturate using 0.2% (w/w) sodium methoxide at 75–80°C for 30 minutes resulted in a $\beta'$-crystallizing fat with good organoleptic properties [64].

List et al. [77] described the preparation of potential margarine and shortening bases by interesterification of vegetable oil and hardstocks (hydrogenated oil or stearin). They found that the interesterified fats possessed plasticity curves similar to those of commercial soft-tub margarine oils prepared by blending hydrogenated hardstocks or commercial all-purpose shortening oils. However, the commercial blends and interesterified blends differed with respect to crystallization behavior.

C. Palm Oil and Palm Kernel Oil

Palm oil has many applications in the food industry. Most often, interesterification of palm oil is combined with hydrogenation and/or fractionation to achieve the most desirable physical and functional properties [14]. Laning [14] described the applications of palm oil in cooking, frying, and salad oils. Corandomization of palm oil with other fats and oils, in combination with fractionation, produced a fluid salad oil.

Cocoa butter, used in the production of chocolate, is expensive and not always available, so substitutes are created, such as those that result from the blending of interesterified lauric acid with other fats. According to Sreenivasan [1], palm kernel oil is a hard butter that melts at 46°C and produces a waxy feel. With interesterification, the melting point is reduced to 35°C. Furthermore, by blending hydrogenated PKO and the randomized product, a whole series of hard butters with highly desirable melting properties is obtained. The effect of randomization on the melting properties of cocoa butter is shown in Figure 15.
D. Milk Fat

Much research has been done on the chemical interesterification of milk fat. Milk fat, like most fats, does not have a random distribution, which conveys a predetermined set of physical properties. Butyric and caproic acids, for example, are predominantly located at sn-3, while palmitic acid is mostly at sn-1 and sn-2 [88]. Other fatty acids are not as specific. Interesterification of milk fat can be a powerful means of modifying its functional properties.

Weihe and Greenbank [89] presented the first paper dealing with the chemical interesterification of milk fat, with details appearing in Weihe [90]. These investigators performed randomization of milk fat at 40–45°C for 20 minutes to 6 hours with 0.1–0.3% Na/K alloy. For directed interesterification, xylene or hexane was added before the reaction, which was begun at 25–38°C and dropped in three to five steps to 10–25°C. Directed interesterification led to more substantial changes than random interesterification (e.g., on solid fat; Fig. 16). Increases in melting point were greater in the presence of a solvent than without, and direct interesterification generated larger increases in melting point than random interesterification.

Interesterification increased the softening point of milk fat by 3.7–4.2°C, which was explained by higher proportion (5–7%) of high-melting triacylglycerols, which translated into a higher hardness. Mickle [91], on the other hand, found that interesterification reduced the hardness of butter and also led to a rancid, metallic flavor. Refining (free fatty acid removal and steam injection under vacuum) removed the undesirable flavor, yet the final product was tasteless. Finally, an in-depth study by Mickle et al. [92] revealed the effects of three interesterification reaction parameters on the hardness of a semisolid resembling butter. All three parameters—duration (5–55 min), temperature (40–90°C), and catalyst concentration (0.5–5%) had sta-
Figure 16  Proportion of solid fat of native butterfat (○), randomized butterfat (●), and butterfat subjected to direct interesterification (□) measured by dilatometry. (Adapted from Ref. 90.)

Statistically significant effects ($p < .05$), with catalyst concentration (at 1–2%) having the greatest influence on hardness, which diminished 45–55%. DeMan [93] observed by means of polarized light microscopy that the crystal habit of interesterified milk fat was markedly changed from that of native milk fat. The effects of cooling procedures on consistency, crystal structure, and solid fat content of milk fat were also examined [94]. Parodi [95] examined the relationship between trisaturates and the softening point of milk fat. Interesteerification increased the softening point from $\sim 32.5^\circ C$ to $\sim 36.5^\circ C$. Timms [68] found that milk fat and beef tallow interesterified blends lacked milk fat flavor. Timms and Parekh [96] explored the possibility of incorporating milk fat into chocolate. Interesteerified milk fat appeared to be better suited to chocolate than noninteresteerified milk fat, but the improvement gained did not compensate for the investment and loss of flavor from interesteerification.

E. Fat Substitutes

Newer applications of chemical interesteerification include the production of low-calorie fat substitutes such as Salatrim and Olestra. Salatrim/Benefat consists of chemically interesteerified mixtures of short chain and long chain fatty acid triacylglycerols. The short chain fraction consists of triacetin, tripropionin, and/or tributyrin, while the long chain fractions consist of hydrogenated soybean oil [26].

Olestra is an acylated sucrose polyester with six to eight fatty acids obtained from vegetable oil (e.g., soybean, corn, sunflower). It is prepared by interesteerifying sucrose and edible oil methyl esters in the presence of an alkali catalyst, at 100–140°C [97]. Olestra is nondigestible, hence noncaloric; it is also nontoxic, yet nutritional concerns potentially exist. Its functionality is dependent on the chain length.
and unsaturation of the esterified fats, as with normal lipids [98]. It can be exchanged for fats in products such as ice cream, margarine, cheese, and baked goods, and it can be blended with vegetable oil.

XI. OXIDATIVE STABILITY

The many advantages of chemical interesterification have been discussed in detail. Many authors have shown, however, that chemical interesterification can negatively influence the oxidative stability of fats and oils. Lau et al. [99] demonstrated that randomized corn oil oxidized three to four times faster than native corn oil. They concluded that the triacylglycerol structure probably was implicated, but the mechanisms remained unclear. Lo and Handel [61] showed that interesterified blends of soybean oil and beef tallow were more unstable following interesterification.

Gavriilidou and Boskou [100] examined the effects of chemical interesterification on the autoxidative stability of an 80% olive oil–20% tristearin blend. The randomized fats were less stable than the native mixtures (Fig. 17). Addition of BHT stabilized the fats, resulting in a peroxide value similar to that for commercially processed hydrogenated vegetable oil used in margarine.

An important contribution to the literature was made by Zalewski and Gaddis [101], who investigated the effect of transesterification of lard on stability, antioxidant efficiency, and rancidity development. Interesteerification of lard did not affect its resistance to oxidation, but changes in oxidative stability due to tocopherol decomposition and the formation of reducing substances were noted. In the absence of antioxidants, both interesterified and native lard had similar peroxide values. Furthermore, because of the position of unsaturated fatty acids at 1,3-positions or randomization toward the 2-position in pork fat triacylglycerols, there was no appreciable effect on initiation of oxidation and autoxidation rates.

Figure 17  Change in peroxide value of an 80%:20% olive oil–tristearin blend before and after interesterification: ●, native blend; ●, randomized blend; □, hydrogenated blend; ■, randomized blend + BHT. (Adapted from Ref. 100.)
Tautorus and McCurdy [102] demonstrated the effects of chemical and enzymatic randomization on the oxidative stability of vegetable oils stored at different temperatures. Noninteresterified and interesterified oils (canola, linseed, soybean, and sunflower) stored at 55°C demonstrated little difference to lipid oxidation, whereas noninteresterified samples were more stable at 28°C. Samples at 55°C underwent much greater oxidation than the samples at 28°C.

Park et al. [103] found that loss of tocopherols accelerated the autoxidation of randomized oils. α-Tocopherol was not detectable following interesterification, while γ-tocopherol and δ-tocopherol diminished 12% and 39%, respectively.

Konishi et al. [104] found that regioselectively interesterified blends of methyl stearate and soybean oil had increased oxidative stability over both native and randomized blends, as monitored by peroxide value and volatiles analysis. The improved oxidative stability was presumably due to the regioselective incorporation of stearic acid at the sn-1(3) carbon sites of the triacylglycerol moiety, which stabilized the linoleic acid, predominantly located at the sn-2 position.

XII. NUTRITIONAL CONSEQUENCES OF INTERESTERIFICATION

Perhaps chemical interesterification’s greatest advantage over hydrogenation lies in nutrition. At present, there are still unsettled nutritional concerns regarding trans fatty acids and their possible links to coronary heart disease [105–107]. Trans fatty acids are present in many edible fats and oils produced worldwide, yet these substances occur in great proportions in partially hydrogenated margarines. Barring non-hydrogenated margarines, literature data indicate that the typical trans fatty acid content of margarines is 10–27% in the United States and 10–50% in Canada [108,109].

It has been shown that randomization does not influence the nutritional value of unsaturated fatty acids [110]. However, not much is known about the potential importance of stereospecificity in the biological activity of dietary fatty acids [111]. In clinical trials, substitution of randomized butter for natural butter tended to reduce serum triacylglycerol and cholesterol concentrations [112]. Human infants absorbed 88% stearic acid when fed lard but only 40% when fed randomized lard. Hence, absorbability and pharmacological properties of fatty acids can be influenced by the molecular form in which they are absorbed [63].

De Schrijver et al. [111] examined lipid metabolism response in rats fed tallow, native or randomized fish oil, and native or randomized peanut oil and found that randomized lard had no significant effects on any of the lipid measurements. Absorption of oleic acid and polyunsaturated fatty acids did not depend on the fatty acid profile of dietary fat. Kritchevsky [113] found that peanut oil’s tendency to produce atherogenicity in rabbits disappeared following chemical interesterification.

It is known that human milk is well absorbed in part because of its proportion of long chain saturated fatty acids located at the sn-2 position. Lien et al. [114] found that mixtures of coconut oil and palm olein were better absorbed by rats if the proportion of long chain saturated fatty acids at the sn-2 position was increased by random chemical interesterification.

Mukherjee and Sengupta [115] found that interesterified soya–butterfat feeding significantly decreased serum cholesterol in humans and rats. The decrease was greater than when noninteresterified blends were fed. The lowering of serum cho-
lesterol paralleled the decrease in concentration of trisaturates and the scattering of 
myristic acid away from sn-2 to sn-1 and sn-3 positions.

Finally, Koga et al. [116] examined the effects of randomization of partially 
hydrogenated corn oil on fatty acid and cholesterol absorption and on tissue lipid 
levels in rats. They found that interesterification did not lead to beneficial effects but 
rather enhanced the hypercholesterolemic tendency of trans fatty acids.

There appears to be some dispute as to the health effects of interesterification. 
Whereas the dietary concerns for avoiding trans fatty acids seem well documented, 
the nutritional effects of fatty acid positional distribution are presently less clear-cut.

XIII. DISTINGUISHING CHEMICAL FROM ENZYMATIC 
INTERESTERIFICATION

Although great strides have been made with extracellular microbial lipases as cata-
lysts for interesterification, most of the industry still relies on chemical interesteri-

cation. Each type of interesterification possesses advantages and disadvantages. Ad-
vantages of chemical interesterification over enzymatic transformations primarily 
involve cost recovery and initial investment. Chemical catalysts are much cheaper 
than lipases. Even with immobilization procedures, capital investment remains high. 
Second, chemical interesterification is a tried-and-true approach; it has been around 
for a long time, and industrial procedures and equipment are available [45].

Costs aside, does treatment by means of chemical or enzymatic interesterifi-
cation in identical applications result in the same final product? Kalo et al. [117] 
compared the changes in triacylglycerol composition and physical properties of but-
terfat interesterified using either sodium methoxide or a nonspecific lipase from Cac-
dida cylindraceae and found only small differences in both interesterified butterfats. 
The compositional changes induced by both chemical and enzymatic means were 
similar, with the trisaturated triacylglycerol content being slightly higher in the en-
zymatically modified product. In terms of physical properties, the chemically inter-
esterified butterfat was slightly harder than its enzymatically modified counterpart. 
Hence, for randomization purposes, the methods appeared to yield similar results for 
the modification of butterfat. However, the product’s butter flavor must be taken into 
account. The harsh process conditions of chemical interesterification result in loss of 
butter’s fine flavor. For purposes where flavor is not a problem, the simpler, tried-
and-true chemical process is preferable.

Enzymatic interesterification has many advantages, such as milder processing 
conditions and the possibility of regiospecificity and fatty acid specificity. This spec-
icity permits structuring not possible by chemical means. For the production of 
nutritionally superior fats, enzymatic interesterification is ideally suited.

XIV. PERSPECTIVES

Chemical interesterification is likely to remain a force in the food industry for the 
foreseeable future. With the progressive demise of hydrogenation likely to continue, 
interesterification (both chemical and enzymatic) will gain greater prominence as a 
food lipid modification strategy.
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REFERENCES


Lipid Oxidation of Edible Oil

DAVID B. MIN and JEFFREY M. BOFF

The Ohio State University, Columbus, Ohio

I. INTRODUCTION

Oxidation can alter the flavor and nutritional quality of foods and produce toxic compounds, all of which can make the foods less acceptable or unacceptable to consumers [1]. Oxidation products typically include low molecular weight compounds that are volatile as well as undesirable off-flavor compounds [2,3]. Triplet oxygen lipid oxidation of foods has been extensively studied during the last 70 years as part of the effort to improve the oxidative stability of foods [4,5]. However, triplet oxygen oxidation does not fully explain the initiation step of lipid oxidation [6,7]. Rawls and VanSanten [8] suggested that singlet oxygen is involved in the initiation of triplet oxygen lipid oxidation because singlet oxygen can directly react with double bonds without the formation of free radicals. Singlet oxygen oxidation can be very rapid in foods due to the low activation energy required for the chemical reaction. The reaction rates of singlet oxygen and triplet oxygen with linoleic acid are $1.3 \times 10^5$ M$^{-1}$ s$^{-1}$ and $8.9 \times 10^4$ M$^{-1}$ s$^{-1}$, respectively [8,9]. For many foods containing lipids, special measures are taken to reduce or prevent oxidation, such as removal of oxygen, addition of antioxidants, and use of gas barrier packaging materials.

Rate of oxidation is dependent on several factors, including temperature, presence of inhibitors or catalysts, and nature of the substrates [10]. Unsaturated fatty acids are more susceptible to oxidation than saturated fatty acids, a property that is primarily due to the lowered activation energy in the initiation of free radical formation for triplet oxygen autoxidation [11,12]. During the last 30 years, increased attention has been given to singlet oxygen oxidation of foods. The significance of singlet oxygen oxidation in foods can be summarized in two statements. (a) The rate of singlet oxygen oxidation is much greater than that of triplet oxygen oxidation,
resulting in drastically increased rates of oxidation even at very low temperatures, lowering the quality of foods during processing and storage [8]. (b) Singlet oxygen oxidation can produce compounds absent in triplet oxygen oxidation due to the different reaction mechanisms of singlet oxygen from the triplet oxygen [6,13].

This chapter reviews the important chemical mechanisms involved in the oxidation of edible oils by singlet and triplet oxygen for the formation of volatile compounds and the effects of the process on the flavor quality of edible oils. Singlet oxygen oxidation will be emphasized in that its importance to lipid oxidation has received increasing attention recently.

II. TRIPLET OXYGEN AND SINGLET OXYGEN CHEMISTRY

Differences in the chemical properties of triplet and singlet oxygen are best illustrated by their molecular orbitals. The molecular orbital of triplet oxygen is shown in Figure 1. The spin multiplicity used to define spin states of molecules is defined as $2S + 1$, where $S$ is the total spin quantum number. The total spin quantum number ($S$) of triplet oxygen is 1. Triplet state oxygen has three closely grouped energy states by two unpaired electrons under a magnetic field. Therefore, the triplet state oxygen has paramagnetic and diradical properties and gives spin multiplicity of 3. Triplet oxygen reacts with radical compounds in foods. Most compounds are in the singlet state, but the most abundant and stable atmospheric oxygen is in triplet state. The molecular orbital of singlet oxygen differs from that of triplet oxygen in that electrons in the antibonding orbital are paired, as shown in Figure 2. The molecule is singlet if the resultant spin ($S$) is zero, dictating the multiplicity of the state, $2S + 1$, to be 1.

Singlet oxygen is an energetic molecule that is in violation of Hund’s rule and the resulting electronic repulsion can produce five excited state conformations. The $1\Delta$ state of singlet oxygen is responsible for most singlet oxygen oxidation in foods and is generally referred to as singlet oxygen [13]. The most energetic electrons of the activated $1\Delta$ state have opposite spins and lie in one single orbital, as shown in Figure 2. Its energy is 22.4 kcal above the ground state of triplet oxygen and exists long enough to react with other singlet state molecules [14]. Singlet oxygen is not a radical compound and can only react with nonradical, singlet state, double-bonded compounds. The lifetime of singlet oxygen is from 50 to 700 $\mu$s, depending on the solvent system of foods [15]. The reaction temperature has little effect on the oxidation rate of singlet oxygen with foods due to the low activation of 0–6 kcal/mol [15]. A summary of chemical properties of singlet and triplet oxygen is shown in Table 1.

III. SINGLET OXYGEN FORMATION

Singlet oxygen can be formed chemically [16], enzymatically [17,18], photochemically [19], and by decomposition of hydroperoxides [20]. Photosensitizers such as chlorophyll, pheophytins, riboflavin, and myoglobin in foods can absorb energy from light and transfer it to triplet oxygen to form singlet oxygen [19,21,22]. Figure 3 shows several pathways for the formation of singlet oxygen in biological systems [23].
Interaction among light, sensitizer, and oxygen is mainly responsible for singlet oxygen formation in foods [13,24]. The chemical mechanism for the formation of singlet oxygen in the presence of sensitizer, light, and triplet oxygen is shown in Figure 4. The ability of photosensitizers to absorb the energy from light and then pass it to triplet oxygen is a convenient method for transferring energy from a light source to form singlet oxygen. The photosensitizer can absorb light very rapidly, in picoseconds, and becomes an unstable, excited, and singlet state molecule (Sen*). The excited singlet photosensitizer will immediately seek to return to ground state in one of three ways: internal conversion; emission of light; or intersystem crossing (ISC), as shown in Figure 4.

Internal conversion involves the transformation from one excited state to another of the same spin state, resulting in the loss of energy as heat. The sensitizer may return to ground state by the emission of fluorescence. The excited sensitiz
Figure 2  Molecular orbital of singlet oxygen [1].

Table 1  Comparison of Singlet and Triplet Oxygen

<table>
<thead>
<tr>
<th>Factor</th>
<th>(^1\text{O}_2)</th>
<th>(^3\text{O}_2)</th>
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<td>Highly electrophilic</td>
</tr>
<tr>
<td>Reaction</td>
<td>Radical compound</td>
<td>Electron-rich compounds</td>
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Data from Ref. 51.
may also undergo an intersystem crossing in which the excited singlet state molecule becomes an excited triplet state molecule. The excited triplet state sensitizer may then react with triplet state atmospheric oxygen to form singlet oxygen and singlet sensitizer via triplet–triplet annihilation mechanism.

IV. TYPE I AND TYPE II PATHWAYS OF EXCITED TRIPLET SENSITIZER

The excited triplet sensitizer may interact directly with substrate such as linoleic acid or phenol compounds by donating an electron or accepting hydrogen, resulting in the production of free radicals or free radical ions as shown in Figure 5. This mechanism is known as type 1 pathway [25,26]. The photosensitizer acts as a photochemically activated free radical initiator. The product radicals have a variety of possible reactions, such as reaction with, or electron transfer to, oxygen and electron or hydrogen abstraction from other substrates. After initiation of the free radicals (R·), the radical compound may react with triplet oxygen via free radical oxidation mechanisms. The rate of the type 1 pathway is mostly dependent on the type and concentration of the sensitizer and substrate. Compounds that are readily oxidizable compounds, such as phenols, and easily reducible compounds, such as quinones, tend to favor type 1 pathway [14].

The excited triplet sensitizer is capable of interacting with triplet oxygen in one of two manners, known as the type 2 pathway. The first, and most significant,
Figure 4  Excitation and deactivation of photosensitizer for the formation of singlet oxygen [13].

involves the generation of singlet oxygen via triplet sensitizer–triplet oxygen annihilation. Upon collision of the excited triplet sensitizer and triplet oxygen, energy is transferred from the sensitizer to the oxygen, resulting in the generation of singlet oxygen and singlet sensitizer. The alternative interaction between triplet sensitizer and triplet oxygen occurs by electron transfer from the excited triplet sensitizer to triplet oxygen, resulting in the formation of superoxide ion. Less than 1% of the interactions of triplet sensitizer and triplet oxygen result in the formation of superoxide ion [27]. The rate of type 2 pathway is mostly dependent on the solubility and concentration of oxygen in the food system. Oxygen is more soluble in nonpolar lipids than it is in water [28]. Therefore, if a sensitizer such as chlorophyll were present in soybean oil, the type 2 pathway would be expected to be active. In contrast, water-based food systems such as milk may tend toward the type 1 pathway due to the reduced availability of oxygen. The shift from type 1 to type 2 or vice versa is dependent on the concentration of oxygen and the types and concentration of substrate.

Types 1 and 2 reactions will enhance oxidation by either the formation of reactive substrate species or the production of singlet oxygen. The competition be-

Figure 5  Formation of excited triplet sensitizer and its reaction with substrate (type I) and triplet oxygen (type II) [13].
tween substrate and triplet oxygen for the excited triplet sensitizer largely determines whether the reaction pathway is type 1 or type 2. Photosensitized oxidation may change the types of pathway during the course reaction as the concentration of substrate and oxygen changes. In aqueous–lipid biphasic systems, the longer half-life of singlet oxygen in the lipid phase favors the oxidation of compounds that partition into the lipids.

V. ELECTRON SPIN RESONANCE SPECTROSCOPY OF SINGLET OXYGEN

The detection of singlet oxygen during photosensitized oxidation is very difficult in that its lifetime is only microseconds. Electron spin resonance (ESR) spectroscopy is a highly sensitive analytical method for the detection of free radicals. ESR detected the formation of singlet oxygen in meat and milk using a spin trapping technique [29,30]. A spin trapping agent, such as 2,2,6,6-tetramethyl-4-piperidone (TMPD), can react with singlet oxygen to form a stable nitroxide radical adduct, 2,2,6,6-tetramethyl-4-piperidone-\(N\)-oxyl (TAN), which is easily measured by ESR. The reaction of TMPD with singlet oxygen for 2,2,6,6-4-piperidone-\(N\)-oxyl formation is shown in Figure 6.

Addition of TMPD to both skim and whole milk during illumination produced TAN after only 5 minutes, confirming the formation of singlet oxygen in milk under light [13]. The limitation of this method is that concentrations of TAN must remain over 10\(^{-8}\) M for detection and over 10\(^{-6}\) M for good spectral resolution. Since the lifetime of singlet oxygen is less than 1 ms, steady-state concentrations greater than 10\(^{-7}\) M are rarely maintained. Coupling spin trapping with ESR spectroscopy could improve this technique of measuring singlet oxygen concentration. Effects of 0, 5 and 15 minutes illumination on ESR spectrum of 2,2,6,6-4-piperidone-\(N\)-oxyl in water solution of riboflavin and 2,2,6,6-tetramethyl-4-piperidone is shown in Figure 7 [5].

VI. REACTIONS OF SINGLET OXYGEN

Singlet oxygen, an electrophilic molecule, is seeking electrons to fill its vacant molecular orbital. One of the most important reaction characteristics of singlet oxygen is that it can directly react with the electron-rich double bonds of unsaturated molecules [31]. The reaction rates of singlet oxygen with oleic, linoleic, linolenic, and arachidonic acids are 0.74, 1.3, 1.9, and 2.4 \(\times\) 10\(^{5}\) M\(^{-1}\) s\(^{-1}\), respectively, which is

![Figure 6](image-url)  
Figure 6 Reaction of 2,2,6,6-tetramethyl-4-piperidone with singlet oxygen for 2,2,6,6-4-piperidone-\(N\)-oxyl formation [30].
relatively proportional to the number of double bonds in the molecules instead of types of double bonds, such as conjugated double bonds [9].

The reaction of linoleic acid with singlet oxygen for conjugated and nonconjugated hydroperoxide formation by ene reaction is shown in Figure 8. Singlet oxygen directly interacts with double bonds to form both conjugated and nonconjugated diene hydroperoxides, a property that is different from triplet oxygen oxidation, which only produces conjugated diene hydroperoxides from linoleic and linolenic acids. Direct interaction of singlet oxygen with double bonds also permits the for-
Figure 8 Conjugated and nonconjugated hydroperoxide formation from linoleic acid by the ene reaction of singlet oxygen [1].

Singlet oxygen participates in reactions such as 1,4-cycloaddition to diene and heterocyclic compounds [31,33], the “ene” reaction [19,34], and 1,2-cycloaddition to olefins [35], all of which involve interaction with double bonds, as shown in Figure 9. Singlet oxygen oxidation of phenols to form dienones via electron transfer and of sulfides to form sulfoxides as shown in Figure 9 is also important in food systems [25].

VII. REACTIONS OF TRIPLET OXYGEN

Diradical triplet oxygen can react with radical food compounds. The initiation of radical formation in the food molecule will be at the site most liable for the loss of a hydrogen atom [10]. The removal of hydrogen from a saturated fatty acid requires approximately 100 kcal/mol of energy. The energy required for the removal of hydrogen at different carbons of linoleic acid is quite different, as shown in Figure 10. A hydrogen at position 11 of linoleic acid is most easily removed due to the presence of a double bond on both sides, requiring only about 50 kcal/mol.

Once the hydrogen is removed, a pentadienyl radical intermediate between carbon 9 and carbon 12 of linoleic acid is formed. The pentadienyl radical provides an equal mixture of conjugated 9- and 13-diene radical and produces 9- and 13-conjugated diene hydroperoxides upon reaction with triplet oxygen, as shown in Figure 11. Triplet oxygen autoxidation results in only the conjugated diene hydroperoxides in linoleic and linolenic acids. The relative reaction ratio of triplet oxygen with oleic, linoleic, and linolenic acid for hydroperoxide formation is 1:12:25, which is dependent on the relative difficulty for the radical formation in the molecule. The reaction rate of triplet oxygen with linolenic acid is about twice as fast as that of linoleic acid because linolenic acid has two pentadienyl groups in the molecule, compared with the linoleic acid with one pentadienyl group.

VIII. COMPARISON OF SINGLET OXYGEN AND TRIPLET OXYGEN

Neff and Frankel [37] compared triplet oxygen and photosensitized singlet oxygen oxidation of oleic, linoleic, and linolenic acids. The hydroperoxides formed between triplet oxygen or singlet oxygen with oleic, linoleic, and linolenic acids are shown
Figure 9  Reactions of singlet oxygen with olefins, sulfides, and phenols [36].

In Table 2, hydroperoxides formed by singlet oxygen oxidation are at positions that formerly contained double bonds. Singlet oxygen produced conjugated and nonconjugated hydroperoxides from linoleic and linolenic acids, but the triplet oxygen produced only conjugated hydroperoxides from linoleic and linolenic acids. The relative reaction rates of triplet oxygen and singlet oxygen with oleic, linoleic, and linolenic

![Chemical structures and reactions](https://example.com/chemistry.png)

Figure 10  Energy required for hydrogen removal from linoleic acid.
**Figure 11**  Conjugated hydroperoxide formation from linoleic acid by free radical reaction of triplet oxygen [1].

<table>
<thead>
<tr>
<th></th>
<th>Oleate</th>
<th>Linoleate</th>
<th>Linolenate</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Singlet oxygen</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conjugated hydroperoxides</td>
<td>9-OOH</td>
<td>9-OOH</td>
<td>9-OOH</td>
</tr>
<tr>
<td></td>
<td>10-OOH</td>
<td>10-OOH</td>
<td>10-OOH</td>
</tr>
<tr>
<td><strong>Nonconjugated hydroperoxides</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>10-OOH</td>
<td>10-OOH</td>
<td>10-OOH</td>
</tr>
<tr>
<td></td>
<td>12-OOH</td>
<td>12-OOH</td>
<td>15-OOH</td>
</tr>
<tr>
<td><strong>Triplet oxygen</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>8-OOH</td>
<td>9-OOH</td>
<td>9-OOH</td>
</tr>
<tr>
<td></td>
<td>9-OOH</td>
<td>10-OOH</td>
<td>12-OOH</td>
</tr>
<tr>
<td></td>
<td>10-OOH</td>
<td>11-OOH</td>
<td>16-OOH</td>
</tr>
<tr>
<td><strong>Conjugated hydroperoxides</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>9-OOH</td>
<td>9-OOH</td>
<td>9-OOH</td>
</tr>
<tr>
<td></td>
<td>13-OOH</td>
<td>12-OOH</td>
<td>13-OOH</td>
</tr>
<tr>
<td></td>
<td>13-OOH</td>
<td>13-OOH</td>
<td>16-OOH</td>
</tr>
</tbody>
</table>

Data from Refs. 1 and 32.
Table 3  Relative Oxidation Rates of Triplet and Singlet Oxygen with Oleate, Linoleate, and Linolenate

<table>
<thead>
<tr>
<th></th>
<th>C18:1</th>
<th>C18:2</th>
<th>C18:3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Triplet oxygen</td>
<td>1</td>
<td>27</td>
<td>77</td>
</tr>
<tr>
<td>Singlet oxygen</td>
<td>$3 \times 10^5$</td>
<td>$4 \times 10^5$</td>
<td>$7 \times 10^4$</td>
</tr>
</tbody>
</table>

*Source: Ref. 38.*

Acids are shown in Table 3. The reaction rates of triplet oxygen and singlet oxygen with linoleic acid are $8.9 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ and $1.3 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$, respectively [8]. That is, single oxygen reacts with linoleic acid about 1450 times faster than the triplet oxygen reacts with linoleic acid.

**IX. DECOMPOSITION OF HYDROPEROXIDES**

The decomposition of hydroperoxides to produce volatile compounds is shown in Figure 12. The most likely decomposition pathway of hydroperoxide is the cleavage

![Figure 12](image_url)

**Figure 12**  Decomposition of hydroperoxides to produce volatile compounds.
between the oxygen and the oxygen of the R-O-O-H, i.e., R-O-O-H R-O + O-H instead of R-O-O-H R-O-O-H. Hiatt and others [39] reported that the activation energy of the cleavage of R-O-O-H was 44 kcal/mol, compared with the cleavage between the oxygen and hydrogen of R-O-O-H, which has high activation energy (90 kcal/mol). Therefore, the hydroperoxide groups are cleaved by homolysis to yield an alkoxy and a hydroxy radical, as shown in Figure 12.

The alkoxy radical formed from hydroperoxide is cleaved by the hemolytic β scission of a carbon–carbon bond to produce oxo compounds and an alkyl or alkenyl radical. The hemolytic β scission is an important free radical reaction that produces volatile compounds in edible oils during oxidation. The unsaturated alkoxy radical can be cleaved by β scission in two mechanisms of cleavage (A and B of Fig. 12). Scission of the carbon–carbon bond on the side of the oxygen-bearing carbon atom will result in formation of unsaturated oxo compounds and an alkyl radical, while scission of the carbon–carbon bond between the double bond and the carbon atom bearing the oxygen will produce a 1-olefin radical and an alkyl oxo compound. The alkyl radical can combine with hydroxy radical to produce an alcohol, and the 1-olefin radical can be converted to a 1-enol. The 1-enol will produce the corresponding oxo compound by tautomerization. An alternative reaction of the radicals eliminated by β scission of the alkoxy radical is hydrogen abstraction from a compound RH. The 8-hydroperoxy methyl oleate can produce 2-undecenal, methyl 7-hydroxyheptanoate, methyl heptanoate, methyl 8-oxo-octanoate, decanal, and 1-decene. The 13-hydroperoxy methyl linoleate can produce methyl 13-oxo-9,11-tridecadienoate, 1-pentanol, pentane, hexanal, methyl 12-oxo-10-dodecenoate, and methyl 9,11-dodecadienoate. Frankel [10] reported the volatile compounds formed from methyl oleate hydroperoxides formed by free radical oxidation and photosensitized oxidation (Table 4).

Frankel [10] reported that the amounts of 2-decenal, 9-oxodecanoate from triplet oxygen oxidation of hydroperoxides differed from those formed in photosensitized singlet oxygen oxidation, as shown in Table 4. Frankel [10] demonstrated that 9- and 10-hydroperoxides formed in the photosensitized oxidation of oleic acid are isomerized into a mixture of 8-, 9-, 10-, and 11-hydroperoxides at 210°C, as shown in Table 4. Chan and others [40] reported that the relative concentrations of hexanal, methyl octanoate, 2,4-decadienal, and methyl 9-oxononanoate from 9-monohydroperoxide of methyl linoleate were 1%, 37%, 51%, and 12%, respectively. However, the relative concentrations of hexanal, methyl octanoate, 2,4-decadienal, and methyl 9-oxononanoate formed from 13-monohydroperoxide of methyl linoleate were 28%, 24%, 33%, and 16%, respectively. There was a significant difference in the composition of volatile compounds formed from 9-monohydroperoxide and 13-monohydroperoxide of methyl linoleate. Different amounts of pentane are formed from linoleic acid by triplet oxygen and singlet oxygen.

X. FLAVOR PROPERTIES OF VOLATILE COMPOUNDS FROM OIL OXIDATION

The types of volatile compounds produced from the oxidation of edible oils are influenced by the composition of the hydroperoxides and the types of oxidative cleavage of double bonds in the fatty acids [41]. A variety of compounds, such as hydrocarbons, alcohols, furans, aldehydes, ketones, and acid compounds, are formed
Table 4 Volatile Compounds Formed from Methyl Oleate Hydroperoxides

<table>
<thead>
<tr>
<th>Compound</th>
<th>Autoxidation (rel. %)</th>
<th>Photosensitized oxidation (rel. %)</th>
<th>Origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heptane</td>
<td>4.4</td>
<td>4.6</td>
<td>11-OOH</td>
</tr>
<tr>
<td>Octane</td>
<td>2.7</td>
<td>10</td>
<td>10-OOH</td>
</tr>
<tr>
<td>Heptanal</td>
<td>0.5</td>
<td>0.5</td>
<td>?</td>
</tr>
<tr>
<td>1-Heptanol</td>
<td>0.4</td>
<td>0.4</td>
<td>11-OOH</td>
</tr>
<tr>
<td>Octanol</td>
<td>11</td>
<td>3.8</td>
<td>11-OOH</td>
</tr>
<tr>
<td>Methyl heptanoate</td>
<td>1.5</td>
<td>4.9</td>
<td>8-OOH</td>
</tr>
<tr>
<td>1-octanol</td>
<td>0.4</td>
<td>1.0</td>
<td>10-OOH</td>
</tr>
<tr>
<td>Nonanal</td>
<td>15</td>
<td>10</td>
<td>9/10-OOH</td>
</tr>
<tr>
<td>Methyl octanoate</td>
<td>5.0</td>
<td>9.7</td>
<td>9-OOH</td>
</tr>
<tr>
<td>2-Nonenal</td>
<td>0.5</td>
<td>0.7</td>
<td>?</td>
</tr>
<tr>
<td>Decenal</td>
<td>3.9</td>
<td>2.0</td>
<td>8-OOH</td>
</tr>
<tr>
<td>Methyl nonanoate</td>
<td>1.5</td>
<td>0.8</td>
<td>?</td>
</tr>
<tr>
<td>2-Decenal</td>
<td>5.4</td>
<td>12</td>
<td>9-OOH</td>
</tr>
<tr>
<td>2-Undecenal</td>
<td>1.7</td>
<td>7.1</td>
<td>8-OOH</td>
</tr>
<tr>
<td>Methyl 8-oxooctanoate</td>
<td>3.5</td>
<td>3.0</td>
<td>8-OOH</td>
</tr>
<tr>
<td>Methyl 9-oxononanoate</td>
<td>15</td>
<td>11</td>
<td>9/10-OOH</td>
</tr>
<tr>
<td>Methyl 10-oxodecanoate</td>
<td>12</td>
<td>1.7</td>
<td>11-OOH</td>
</tr>
<tr>
<td>Methyl 10-oxo-8-decanoate</td>
<td>3.4</td>
<td>5.0</td>
<td>10-OOH</td>
</tr>
</tbody>
</table>

Source: Ref. 10.

during oxidation. Most of these are responsible for the off-flavor in oxidized edible oils. However, the aliphatic carbonyl compounds, such as alkanals, trans,trans-2,4-alkadienals, isolated alkadienals, isolated cis-alkenals, trans,cis-2,4-alkadienals, and vinyl ketones, have the lowest threshold values as shown in Table 5. Frankel [10] reported that the significant compounds responsible for flavor are trans,cis-2,4-de-

Table 5 Threshold Values of Compounds Formed from Oxidized Oils

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Threshold (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrocarbons</td>
<td>90–2150</td>
</tr>
<tr>
<td>Substituted furans</td>
<td>2–27</td>
</tr>
<tr>
<td>Vinyl alcohols</td>
<td>0.5–3</td>
</tr>
<tr>
<td>1-Alkenes</td>
<td>0.02–9</td>
</tr>
<tr>
<td>2-Alkenals</td>
<td>0.04–2.5</td>
</tr>
<tr>
<td>Alkanals</td>
<td>0.04–1</td>
</tr>
<tr>
<td>trans,trans-2,4-Alkadienals</td>
<td>0.04–0.3</td>
</tr>
<tr>
<td>Isolated alkadienals</td>
<td>0.002–0.3</td>
</tr>
<tr>
<td>Isolated cis-alkenals</td>
<td>0.0003–0.1</td>
</tr>
<tr>
<td>trans,cis-Alkadienals</td>
<td>0.002–0.006</td>
</tr>
<tr>
<td>Vinyl ketones</td>
<td>0.00002–0.007</td>
</tr>
</tbody>
</table>

Source: Ref. 10.
Table 6  Flavor Perceptions of Volatile Compounds Formed by Lipid Oxidation

<table>
<thead>
<tr>
<th>Flavor perception</th>
<th>Responsible compounds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardboard</td>
<td>trans,trans-2,6-Nonadienal</td>
</tr>
<tr>
<td>Oily</td>
<td>Aldehydes</td>
</tr>
<tr>
<td>Painty</td>
<td>Pent-2-enal, aldehydes</td>
</tr>
<tr>
<td>Fishy</td>
<td>trans,trans-2,4,7-Decatrienol, oct-1-en-3-one</td>
</tr>
<tr>
<td>Grassy</td>
<td>trans-2-Hexenal, nona-2,6-dienal</td>
</tr>
<tr>
<td>Deep-fried</td>
<td>trans,trans-2,4-Decadienal</td>
</tr>
</tbody>
</table>

Source: Ref. 41.
ganoleptic evaluation of the addition of 2 ppm 2-pentylfuran to freshly deodorized and bland soybean oil was found to provide the “reverted” soybean oil flavor. The addition of 2 ppm 2-pentylfuran to deodorized cottonseed oil and corn oil produced reversion flavor found in reverted soybean oil [48].

Ho and others [46] determined that 2-(1-penteny)lfuran contributed to reversion flavor. In a similar study, Smagula and others [49] reported that 2-(2-penteny)lfuran is also a contributor according to detailed organoleptic evaluation. Flavor thresholds of the 2-pentenylfuran isomers were found to be between 0.25 and 6 ppm. Chang and others [50] isolated and identified all four 2-pentenylfuran isomers in reverted soybean oil.

Mechanisms for the formation of 2-pentylfuran from linoleic acid and 2-pen-tenylfuran isomers from linolenic acid using triplet oxygen were proposed by Chang and others [48] and Ho and others [46], respectively. Formation of both 2-pentylfuran and 2-pentenylfuran requires a hydroperoxide intermediate at carbon 10. Formation of 10-hydroperoxide in linoleic or linolenic acids by free radical triplet oxygen oxidation is highly improbable, but is very common in the singlet oxygen oxidation of linoleic or linolenic acids as shown in Table 2. Min [51] reported that the chemical mechanisms for the formations of 2-pentylfuran from linoleic acid and 2-(2-pentenyl)lfuran formation from linolenic acid using singlet oxygen as shown in Figures 14 and 15, respectively.

Callison [52] identified 2-pentylfuran and 2-pentenylfuran in soybean oil containing 5 ppm chlorophyll $b$ during storage under light for 96 hours, as shown in

**Figure 13** The effects of 0, 2, 4, 6, and 8 ppm chlorophyll on the formation of head space volatile compounds of soybean oil under light [44].
Figure 14. Mechanism for the formation of 2-pentylfuran from linoleic acid by singlet oxygen [51].

Figure 16. 2-Pentylfuran was identified by comparing the gas chromatographic retention time to that of standard 2-Pentylfuran compound. 2-Pentenylfuran was identified by mass spectrum, as shown in Figure 17. The formation of 2-pentenylfuran increased with increasing storage time and concentration of chlorophyll, as shown in Table 7. 2-Pentylfuran and 2-(2-pentenyl)furan were formed only in the presence of light and chlorophyll in soybean oil. Soybean oil containing 5 ppm
Figure 15  Mechanism for the formation of 2-pentenylfuran from linolenic acid in soybean oil [51].

chlorophyll did not produce 2-pentylfuran and 2-pentenyl furan during dark storage. The soybean oil with no chlorophyll (removed by silicic acid chromatography) did not produce 2-pentylfuran or 2-pentenylfuran during light storage. These results clearly suggest the importance of light and chlorophyll in the stability of soybean oil and support the singlet oxygen oxidation theory of the formation of 2-pentylfuran and 2-pentenylfuran, which are contributors to reversion flavor.
Figure 16  Gas chromatogram of soybean oil with 5 ppm chlorophyll added (a) initially and (b) after storage under light for 4 days [52].
XII. SINGLET OXYGEN QUenchING MECHANISMS

Other than exclusion of light and reduction of oxygen present, the use of quenching agents is perhaps the best way to reduce singlet oxygen oxidation and may be the only protective measure if the other two alternatives are not feasible. Natural food components, such as tocopherols, carotenoids, and ascorbic acid, can act as effective quenching agents [53]. The quenching agent may interfere with the development or activity of singlet oxygen at several stages of development. Figure 18 shows the development of singlet oxygen and its subsequent reaction with substrate (A) to form the oxidized product (AO₂).

At every stage in this reaction there is at least one alternate route, which, if taken, would minimize the oxidation of the substrate (A). The first step represents the return of the excited singlet sensitizer (¹Sen*) to ground state (¹Sen) without ISC to form the excited triplet sensitizer (³Sen*). The second represents interaction with a quenching agent (Q) at a rate represented as k_Q, returning the excited triplet sensitizer (³Sen*) to ground state (¹Sen) prior to interaction with triplet oxygen. The excited triplet sensitizer (³Sen*) may interact with triplet oxygen (O₂) to form singlet oxygen (¹O₂). Following its creation, there are three fates for singlet oxygen in foods. (a) It may naturally decay to the ground state at a rate represented as k_d. (b) It may

Table 7 2-Pentenylfuran Peak Areas

<table>
<thead>
<tr>
<th>Light exposure (days)</th>
<th>Added chlorophylla</th>
<th>0 ppm</th>
<th>1 ppm</th>
<th>5 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td></td>
<td>0</td>
<td>0</td>
<td>1502</td>
</tr>
<tr>
<td>4</td>
<td></td>
<td>0</td>
<td>1534</td>
<td>3018</td>
</tr>
</tbody>
</table>

"Electronic counts of gas chromatogram.
Source: Ref. 50."
react with a singlet state substrate (A) at a state represented as \( k_r \), forming the oxidized product \( \text{AO}_2 \). 

(c) It may be destroyed by a quenching agent by either combining with the quencher, at a rate represented as \( k_{\text{Q}} \), to form the product \( \text{QO}_2 \) or by passing its energy to the quenching agent and returning to free triplet oxygen, at a rate represented as \( k_{\text{i}} \).

As represented by Figure 18, there are three points at which a quenching agent may act; one is quenching of the excited triplet sensitizer, and the other two are quenching of singlet oxygen by chemical or physical means. Chemical quenching involves reaction of singlet oxygen with the quenching agent to produce an oxidized product \( \text{QO}_2 \). Physical quenching results in the return of singlet oxygen to triplet oxygen without the consumption of oxygen or product formation achieved by either energy transfer or charge transfer. Therefore, triplet oxygen quenchers must either be able to donate electrons or to accept energy 22.4 kcal above ground state. An example of the latter is \( \beta \)-carotene, which has a low singlet energy state and can therefore accept the energy from singlet oxygen [7]. Ascorbic acid is an example of a chemical that can quench the excited sensitizer. Table 8 lists quenching rates of several quenching agents.

### XIII. CAROTENOIDs

\( \beta \)-Carotene is considered to be the most powerful physical quenching agent in foods. Foote [25] found that one molecule of \( \beta \)-carotene can quench 250–1000 molecules of singlet oxygen at a rate of \( 1.3 \times 10^{10} \, \text{M}^{-1} \text{s}^{-1} \). Energy transfer from singlet oxygen to a quencher results in the formation of triplet oxygen and an excited triplet state quencher. The triplet state quencher may also be formed in the energy transfer reaction between singlet state quencher and an excited triplet state sensitizer in what is referred to as a triplet sensitizer quenching reaction. Investigations into the ability of \( \beta \)-carotene to act as a quencher in food systems has been prominent in recent years.

The rate of singlet oxygen quenching by carotene is dependent on the number of conjugate double bonds in the carotenoid. Lee and Min [7] evaluated the effectiveness of five carotenoids in quenching chlorophyll-sensitized photooxidation of soybean oil and reported that the effectiveness increased with the number of double bonds in the carotenoid and the amount of carotenoid added. Those carotenoids with seven or less double bonds are ineffective as quenchers, being unable to accept the energy from singlet oxygen.
Table 8  Singlet Oxygen Quenchers and Their Quenching Rates

<table>
<thead>
<tr>
<th>Quenching compound</th>
<th>Quenching rate (M⁻¹ s⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>β-Apo-8’-carotenal</td>
<td>3.1 × 10⁹</td>
</tr>
<tr>
<td>β-Carotene</td>
<td>4.6 × 10⁹</td>
</tr>
<tr>
<td>Lutein</td>
<td>5.7 × 10⁹</td>
</tr>
<tr>
<td>Zeaxanthin</td>
<td>6.8 × 10⁹</td>
</tr>
<tr>
<td>Lycopene</td>
<td>6.9 × 10⁹</td>
</tr>
<tr>
<td>Isozeaxanthin</td>
<td>7.4 × 10⁹</td>
</tr>
<tr>
<td>Astaxanthin</td>
<td>9.9 × 10⁹</td>
</tr>
<tr>
<td>Canthaxanthin</td>
<td>11.2 × 10⁹</td>
</tr>
<tr>
<td>α-Tocopherol</td>
<td>2.7 × 10⁷</td>
</tr>
<tr>
<td>1,4-Diazabicyclo-(2,2,2)-octane</td>
<td>1.5 × 10⁷</td>
</tr>
<tr>
<td>Dimethylfuran</td>
<td>2.6 × 10⁷</td>
</tr>
<tr>
<td>BTC⁺</td>
<td>1.2 × 10⁹</td>
</tr>
<tr>
<td>TPB⁺</td>
<td>3.7 × 10⁷</td>
</tr>
</tbody>
</table>

⁺Bis(di-n-butylthiocarbamato)nickel chelate.
⁺[2,2’-Thiobis(4-1,1,3,3-tetramethylbutyl)phenalato]-n-butylamine)
nickel chelate.

Source: Ref. 1.

The quenching ability of β-carotene is the best studied energy-transfer quenching mechanism; however, other compounds with similar properties may be effective singlet oxygen quenchers if their singlet energy level is below 22 kcal. Lee and Min [42] tested the effect of various amounts of β-carotene added to soybean oil that contained 4 ppm chlorophyll and was exposed to light. Figure 19 shows the ability of increasing amounts of β-carotene to reduce the head space oxygen depletion.

XIV. TOCOPHEROLS

Tocopherols are well-studied free radical scavengers. They are the most abundant antioxidants in nature and are the primary antioxidant in vegetable oils. When present in systems that are vulnerable to singlet oxygen oxidation, tocopherols have demonstrated the ability to inhibit lipid peroxidation. Tocopherols were identified in soybean oil averaging about 1100 ppm and exist in α-, β-, γ-, and δ-tocopherol forms at approximately 4%, 1%, 67%, and 29%, respectively. Jung and others [54] studied the effectiveness of α-, γ-, and δ-tocopherol in quenching the photooxidized singlet oxygen oxidation of soybean oil and determined that α-tocopherol had the greatest quenching rate. Foote and others [55] found that the mechanism by which tocopherols act in quenching singlet oxygen involves charge transfer. The reaction involves an electron donation from tocopherol to singlet oxygen, forming a charge transfer complex. The transfer complex undergoes an ISC to ultimately form triplet oxygen and the starting tocopherol. The overall rate is dependent on the ability of the donor to accept electrons in that the most easily oxidized donors are the most efficient quenchers.

Singlet oxygen destruction of vitamin D₂ in a model system was reduced by the presence of α-tocopherol proportional to its concentration in solution [56].
Figure 19  Effect of 0, 0.25, 0.5, and $1.0 \times 10^{-5}$ M $\beta$-carotene on the peroxide value of soybean oil [57].

20 shows the ability of increasing amounts of $\alpha$-tocopherol to reduce the head space oxygen depletion in a model system. The rate of singlet oxygen quenching by $\alpha$-tocopherol is similar to that of $\beta$-carotene.

XV. DETERMINING QUENCHING MECHANISMS

The quenching mechanism of photosensitized singlet oxygen oxidation can be determined by measuring the rate constant of total quenching, physical quenching, and chemical quenching. Quenching agents work in numerous ways to inhibit the formation of oxidized products, as has been previously described (Fig. 18).

The quantum yield of a photochemical reaction is defined as the ratio of the number of molecules of a product formed to the number of photons of light absorbed. This value is used to measure the relative efficiency of a photochemical reaction. The quantum yield of oxidized product formation ($\phi_{AO_2}$) can be defined by the equation:

$$\phi_{AO_2} = A \times B \times C$$  

where A and B represent the partitioning of singlet sensitizer and triplet sensitizer.
toward singlet oxygen formation, respectively, and C represents the formation of the oxidized product.

The amount of quencher necessary to inhibit a substantial amount of the singlet oxygen sensitizer is particularly high and the lifetime of the singlet oxygen sensitizer is very short. For these reasons, singlet sensitizer quenching is not considered in the steady state equation. Therefore, A is a constant (K) that is equal to the quantum yield of ISC.

Term B represents the rate of singlet oxygen formation, which is dependent on the triplet sensitizer quenching rate and the rate of triplet–triplet sensitizer annihilation. Therefore:

\[ B = \frac{k_0 \text{ [oxygen]}}{k_0 \text{ [oxygen]} + k_Q \text{ [quencher]}} \]  

where \( k_0 \) is the reaction rate constant of triplet–triplet annihilation and \( k_Q \) is the reaction rate constant of triplet sensitizer quenching.

Term C represents the formation of oxidized product, which is dependent on the concentration and nature of the substrate, physical and chemical quenching of singlet oxygen, as well as the natural decay rate of singlet oxygen. The assemblage of these factors generates the following equation:
Figure 21  Singlet oxygen quenching mechanism [58].

\[
C = \frac{k_r \text{ [substrate]}}{k_r \text{ [substrate]} + (k_{\alpha\to\gamma} + k_d) \text{ [chemical + physical quencher]} + k_d} \tag{3}
\]

where \( k_r \) is the reaction rate constant of the reaction of singlet oxygen with the substrate, \( k_{\alpha\to\gamma} \) is the reaction rate constant of chemical quenching, \( k_d \) is the reaction rate constant of physical quenching, and \( k_d \) is the decay constant of singlet oxygen.

If a given quenching agent were to inhibit photosensitized oxidation by quenching singlet oxygen, the steady state equation can be written as:

\[
\phi_{\text{AO}} = K \frac{k_r [^3\text{O}_2]}{k_r [^3\text{O}_2] + k_o [Q]} \times \frac{k_r [A]}{k_r [A] + (k_{\alpha\to\gamma} + k_d) [Q] + k_d} \tag{4}
\]

where \( K \) is the quantum yield of ISC of the excited state of the singlet sensitizer [term A from Eq. (1)] and both B and C have been appropriately substituted with Eqs. (2) and (3), respectively.

Figure 22  Triplet sensitizer quenching mechanism [58].
In a given system, if there is only singlet oxygen quenching such that \( k_d[Q] \ll k_i[^3O_2] \), then the B term is equal to 1. Therefore, the steady state equation becomes:

\[
\phi_{AO} = K \frac{k_r[A]}{k_i[A] + (k_{ox-Q} + k_o)[Q] + k_d}
\]  

(5)

This equation can be inverted to

\[
\phi_{AO}^{-1} = K^{-1} \left( 1 + \frac{(k_{ox-Q} + k_o)[Q] + k_d}{k_r[A]} \right)^{-1}
\]  

(6)

so that it is in slope-intercept form.

Alternatively, if there is only triplet sensitizer quenching such that \( (k_{ox-Q} + k_o)[Q] \ll k_i[A] + k_d \), then the slope intercept form of the equation is

\[
\phi_{AO}^{-1} = K^{-1} \left( 1 + \frac{k_o[Q]}{k_i[^3O_2]} \right) \left( 1 + \frac{k_d}{k_i[A]} \right)
\]  

(7)

The significance of these two equations is the fact that one describes a system in which singlet oxygen quenching is dominant and the other describes a system in which triplet sensitizer is dominant. A plot of \([AO_2]^{-1}\) vs. \([A]^{-1}\) at different \([Q]\) will appear in one of two manners, depending on which mechanism dominates a system. If singlet oxygen quenching is dominant [Eq. (6)], then the plots at various \([Q]\) will all have the same \(y\) intercept but different slopes (Fig. 21). If triplet sensitizer quenching is dominant [Eq. (7)], then both the intercept and the slope will vary (Fig. 22).

REFERENCES

58. T. L. Li. Stability and photochemistry of vitamin D₂ in model systems and quenching mechanisms of carotenoids in singlet oxygen oxidation of vitamin D₂. M.S. Thesis. The Ohio State University, Columbus, 1997.
I. INTRODUCTION

Lipid oxidation is one of the major causes of quality deterioration in muscle foods following storage at refrigerated or frozen temperatures. Often seen in later stages of storage, quality losses are manifested through a variety of mechanisms, which are summarized in Table 1 [1–17]. Although lipid oxidation usually causes a decrease in consumer acceptability, in some cases lipid oxidation leads to enhancement of product quality. An example is the enzymatic production of fresh-fish aromas. This chapter reviews the fundamental mechanisms of lipid oxidation as they apply to muscle foods. Included in this review is a discussion of the impact of tissue structure and compositional factors on pathways, kinetics, and extent of oxidation. Also included is a section describing the effect of various food processing applications on lipid oxidation reactions. Throughout this chapter, the reader will be made aware of the multiple interactions among muscle constituents during the process of lipid oxidation. Therefore, a short review (Sec. III.G) details how mathematical models may be used to account for these interactions and indicates how shelf life predictions and conditions for optimal stability may be derived.

II. BASIC CHEMISTRY OF LIPID OXIDATION

The two major components involved in lipid oxidation are unsaturated fatty acids and oxygen. In this process, oxygen from the atmosphere is added to certain fatty acids, creating unstable intermediates that eventually break down to form unpleasant flavor and aroma compounds. Although enzymatic and photogenic oxidation may play a role, the most common and important process by which unsaturated fatty
Table 1  Consequences of Lipid Oxidation Activity

<table>
<thead>
<tr>
<th>Consequence</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh flavors</td>
<td>1,2</td>
</tr>
<tr>
<td>Off-flavors (warmed over/rancid)</td>
<td>3,4</td>
</tr>
<tr>
<td>Cholesterol oxidation products with potentially detrimental health implications</td>
<td>5–9</td>
</tr>
<tr>
<td>Protein denaturation and functionality changes</td>
<td>10–12</td>
</tr>
<tr>
<td>Pigment changes</td>
<td></td>
</tr>
<tr>
<td>Myoglobin (red) → metmyoglobin (brown)</td>
<td>13–15</td>
</tr>
<tr>
<td>Loss of red (carotenoid) pigmentation</td>
<td>16,17</td>
</tr>
</tbody>
</table>

Acids and oxygen interact is a free radical mechanism characterized by three main phases:

Initiation: \[ \text{In}^\cdot + \text{RH} \rightarrow \text{InH} + \text{R}^\cdot \]

Propagation: \[ \text{R}^\cdot + \text{O}_2 \rightarrow \text{ROO}^\cdot \]
\[ \text{ROO}^\cdot + \text{RH} \rightarrow \text{R}^\cdot + \text{ROOH} \]

Termination: \[ 2\text{RO}_2^\cdot \rightarrow \text{O}_2 + \text{RO}_2\text{R} \]
\[ \text{RO}_2^\cdot + \text{R}^\cdot \rightarrow \text{RO}_2\text{R} \]

Initiation occurs as hydrogen is abstracted from an unsaturated fatty acid, resulting in a lipid free radical, which in turn reacts with molecular oxygen to form a lipid peroxy radical. While irradiation can directly abstract this hydrogen from lipids, initiation is frequently attributed in most foods, including muscle foods, to reaction of the fatty acids with active oxygen species. The propagation phase of oxidation is fostered by lipid–lipid interactions, whereby the lipid peroxy radical abstracts hydrogen from an adjacent molecule, resulting in a lipid hydroperoxide and a new lipid free radical. Interactions of this type continue 10 \[18\] to 100 times \[19\] before two free radicals combine to terminate the process. Additional magnification of lipid oxidation, however, occurs through branching reactions (also known as secondary initiation): \[ \text{Fe}^{2+} + \text{LOOH} \rightarrow \text{LO}^\cdot + \text{OH}^\cdot \]. The radicals produced will then proceed to abstract hydrogens from unsaturated fatty acids. Additional information describing these free radical processes is presented later in Sec. I.A, I.B, and I.C, on initiation, propagation, and termination, respectively.

By themselves, lipid hydroperoxides are not considered harmful to food quality; however, they are further degraded into compounds that are responsible for off-flavors. The main mechanism for the formation of aldehydes from lipid hydroperoxides is homolytic scission (\(\beta\) cleavage) of the two C–C bonds on either side of the hydroperoxy group \[20\]. This reaction proceeds via the lipid alkoxyl radical, with the two odd electrons produced on neighboring atoms forming a carbonyl double bond. Two types of aldehydes are formed from the cleavage of the carbon bond: aliphatic aldehydes derived from the methyl terminus of the fatty acid chain and aldehydes still bound to the parent lipid molecule. Since unsaturated aldehydes can be oxidized further, additional volatile products may be formed \[20\].
A. Initiation

The direct reaction of a lipid molecule with a molecule of oxygen is highly improbable because the lipid molecule is in a singlet electronic state and the oxygen molecule has a triplet ground state. To circumvent this spin restriction, oxygen can be activated by any of the following three initiation mechanisms: (1) formation of singlet oxygen; (2) formation of partially reduced or activated oxygen species such as hydrogen peroxide, superoxide anion, or hydroxyl radical; and/or (3) formation of active oxygen–iron complexes (ferryl iron or ferric–oxygen–ferrous complex). In addition, the oxidation of fatty acids may occur either directly or indirectly through the action of enzyme systems, of which three major groups are involved: microsomal enzymes, peroxidases, and dioxygenases, such as lipoxygenase or cyclooxygenase. That such chemical and enzymatic reactions exist in living tissue is evidenced by the occurrence in aerobic organisms of enzymes that can eliminate or detoxify these compounds (e.g., superoxide dismutase, catalase, glutathione peroxidase). Therefore, activated oxygen species are likely to be present in the food item even before it is harvested, not just produced during processing and storage. As for which mechanism of initiation is primarily responsible, a large volume of research has been published exploring this issue but no consensus has arisen. The reader is therefore encouraged to look in the reviews of Kanner et al. [21], Hsieh and Kinsella [22], Kappus [23], and Bradley and Min [24] for a more in-depth look at mechanisms of initiation. However, additional information on specific sources of initiation will be presented in Section III.E.

B. Propagation

Propagation reactions form the basis of the chain reaction process and in general include the following:

Radical coupling with oxygen: \( R' + O_2 \rightarrow ROO' \)

Atom or group transfer: \( ROO' + RH \rightarrow ROOH + R' \)

Fragmentation: \( ROO' \rightarrow R' + O_2 \)

Rearrangement:

Cyclization:

In oxygen radical coupling, molecular oxygen reacts with the carbon-centered free radical at or near the diffusion-controlled rate of approximately \( 10^9 \text{ M}^{-1} \text{ s}^{-1} \). A major consequence of this reactivity is that the concentration of \( R' \) is much smaller than that of \( ROO' \). In atom transfer, whereas a peroxy radical, \( ROO' \), will not readily abstract hydrogen from a saturated hydrocarbon, it will do so very readily from allylic and bisallylic \( \text{C–H} \) bonds of unsaturated fatty acids.
Lower bond energies for bisallylic and allylic hydrogens versus methylene hydrogens (75 and 88 vs. 100 kcal/mol, respectively), as well as resonance stabilization of the radical intermediate, contributes to ease of abstraction from unsaturated fatty acids [25,26]. The newly formed hydroperoxy radical can in turn abstract hydrogen from an adjacent unsaturated fatty acid such that the reaction sequence goes through 8 to 14 propagation cycles before termination [27]. Conditions that determine the chain propagation length include initiation rate, structures of aggregates (increasing with increasing structure of the aggregates), temperature, presence of antioxidants, and chain branching. Chain branching involves the breakdown of fatty acid hydroperoxides to the lipid peroxy or alkoxyl radical. Given the bond dissociation energies of LOO–H (about 90 kcal/mol) and LO–OH (about 44 kcal/mol), spontaneous decomposition is unlikely at refrigerated or freezing temperatures [28]. Instead, breakdown of hydroperoxides would be dominated by one-electron transfers from metal ions during low temperature storage.

\[
Fe^{2+} + LOOH \rightarrow Fe^{3+} + LO^- + OH^-
\]

The major contributors to decomposition of lipid hydroperoxides in food and biological systems would be heme and nonheme iron, with reactions involving the ferrous ion occurring much more quickly than those involving ferric ion.

### III. MUSCLE COMPOSITION AND LIPID OXIDATION

#### A. Muscle Structure and Function

Before discussing individual constituents of muscle tissue, a review of the structural and chemical features that contribute to muscle tissue's oxidative stability will be taken. Within the animal body, there are more than 600 muscles varying widely in shape, size, and activity. However, at the cellular level there is close resemblance among muscles from a wide variety of organisms. The typical arrangement of a skeletal muscle in cross section consists of epimysium (connective tissue surrounding the entire muscle), perimysium (connective tissue separating the groups of fibers into
bundles), and endomysia (sheaths of connective tissue surrounding each muscle fiber). Also surrounding each muscle fiber is the sarcolemma membrane, which periodically, along the length of the fiber, forms invaginations usually referred to as T tubules.

Within the muscle fiber, the sarcoplasm serves to suspend organelles such as mitochondria and lysosomes. Water constitutes 75–80% of the sarcoplasm but in addition may contain lipid droplets, variable quantities of glycogen granules, ribosomes, proteins, nonprotein nitrogenous compounds, and a number of inorganic constituents.

Myofibrils, which are organelles found only in muscle cells, are long, thin, cylindrical rods that extend the entire length of the muscle fiber and constitute the contractile apparatus, which is composed of primarily myosin and actin proteins. Also intracellular in nature is the sarcoplasmic reticulum (SR). Forming a closely meshed network around each myofibril, SR membranes serve as the storage site for Ca$^{2+}$ in resting muscle fibers.

The contractile process is a complex mechanism that starts with an action potential spreading through the T tubules and sarcolemma, finally reaching the sarcoplasmic reticulum. At the SR membrane, depolarization occurs, stimulating the release of calcium. The released calcium binds to troponin causing a conformational change in the protein, which in turn triggers myosin ATPase to hydrolyze ATP to ADP. The chemical energy released from the hydrolysis is utilized by the myosin and actin to initiate the sliding mechanism resulting in the contraction of the muscle. When the impulse that started the potential subsides, the sarcolemma polarizes, stimulating the energy-dependent sequestration of calcium by the SR membranes.

B. Biochemical Changes in Muscle Postmortem

After death of an animal, all circulation ceases—an event that rapidly brings about important changes in the muscle tissue (Table 2). The principal changes are attributable to a lack of oxygen (anaerobic conditions) and the accumulation of certain waste products, especially lactate and H$^+$. In a short time, the mitochondrial system ceases to function in all but surface cells because internal oxygen is rapidly depleted. Anaerobic glycolysis continues to regenerate ATP and lactate, but eventually the decrease in pH caused by the presence of lactate disrupts glycolytic activity. When ATP is depleted, the most immediate response seen is the onset of rigor whereby actin and myosin remain in a contracted state as a result of the absence of a plasticizing agent (ADP or ATP). A more important response with regard to lipid oxidation is the cellular membranes’ inability to maintain their integrity. Consequently, lysosomal enzymes, such as phospholipase and lipase, may be released, affecting in turn the susceptibility of lipids to oxidize. Calcium leakage is a noted response to the increased membrane permeability of SR and mitochondria [30,31]. Increased calcium concentrations, in turn, could activate enzymic systems, such as phospholipase, which would not normally be turned on.

C. Variability Between and Within Muscles

Although the muscle function of locomotion is similar throughout the animal kingdom, compositional differences exist between species and even within different muscles of the same species. Red and white muscles present a classic example, with
**Table 2**  Biochemical Changes and Negative Consequences in Postmortem Muscle Food

<table>
<thead>
<tr>
<th>Biochemical change</th>
<th>Consequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Decrease in ATP</td>
<td>Loss of energy source needed for reduction of many compounds</td>
</tr>
<tr>
<td>Increase in hypoxanthine</td>
<td>Off-flavor produced by ATP degradation</td>
</tr>
<tr>
<td>Conversion of hypoxanthine</td>
<td>Xanthine oxidase can initiate oxidation when molecular oxygen is present in system</td>
</tr>
<tr>
<td>Loss of secondary antioxidants and cofactors</td>
<td>Initiation of reaction</td>
</tr>
<tr>
<td>Decrease in ascorbate, glutathione</td>
<td>Loss of secondary antioxidants and cofactors</td>
</tr>
<tr>
<td>Increase in low molecular weight iron</td>
<td>Initiation of oxidation</td>
</tr>
<tr>
<td>Oxidation of myoglobin to porphyrin radical</td>
<td>Can react with hydrogen peroxide to produce ferryl oxene (Fe⁴⁺), which can initiate lipid oxidation</td>
</tr>
<tr>
<td>Loss of tocopherol</td>
<td>Loss of primary antioxidant</td>
</tr>
<tr>
<td>Disintegration of membranes</td>
<td>Could cause hydrolysis of phospholipids, uneven maintenance of ions</td>
</tr>
<tr>
<td>Loss of Ca²⁺ sequestration</td>
<td>Increased calcium ion content in aqueous phase, causes many inactive processes to become active</td>
</tr>
</tbody>
</table>

White muscles (i.e., muscles dominated by white fibers) presumably best suited for vigorous activity for a short period and red muscles for sustained activity. Red fibers tend to be smaller than white fibers and contain more mitochondria, possess greater concentrations of myoglobin and lipid, have a thicker sarcolemma and a much less extensive and more poorly developed SR, and have less sarcoplasm; however, red fibers are more generously supplied with blood than are white fibers. All of these differences lead to differences in oxidative stability.

Fish muscle is noticeably different from avian or mammalian muscles. Containing a larger percentage of myofibrillar protein than mammalian skeletal muscle, fish are characterized by a large percentage of unsaturated fatty acids. Changes in composition in response to environment are common. For example, fish that live in a low temperature environment have a larger fraction of dark muscle than fish that live in a warmer environment [32]. Caution is necessary when sampling fish for oxidative stability as local differences exist [32–35]. The most susceptible portion of herring fillets to oxidation during ice and frozen storage was found under the skin and was attributed to the large proportion of dark muscle in the sample compared with the middle and inner parts [33,34]. However, isolation and subsequent analysis of white and dark muscle will not always ensure a uniform response to oxidation. While white muscle is considered to be very uniform in composition no matter where it is located on the fish, dark muscle varies in composition as a function of its location, containing more lipid in the anterior part of the fish and more water and protein in the posterior part [32].

To circumvent the inherent variability in composition that occurs between meat cuts, experimental studies often isolate subcellular membranes from muscles and measure their response to oxidative catalysts. Depending on the degree of purification, wide variability may exist among isolated membranes. For instance, a microsomal fraction may contain membranes from SR, mitochondria, Golgi bodies, ly-
sosomes, etc. A comparison of the lipid content of several of these subcellular membranes (Table 3) shows clearly that as the percentage of one membrane type changes in the isolate, the batch lipid composition also changes [36]. Even within one membrane type, such as SR, fractions isolated from cisternal and longitudinal SR [37] have revealed different ratios of phospholipid to protein [38]. Here also the lipid composition of the isolated membranes changes if more of one fraction is isolated. Ultimately, these changes in lipid content will lead to differences in the oxidative susceptibility of the membrane isolate. The next section reviews the effects of lipid composition on oxidative stability.

D. Lipid Substrate

1. Fatty Acid Unsaturation

Carbon–hydrogen bond dissociation energies of a fatty acid are lowest at bisallylic methylene positions. These are the positions between adjacent double bonds [26,39]. Consequently, these positions are the thermodynamically favored sites for attack by lipid peroxyl radicals in polyunsaturated fatty acids (PUFAs). In studies involving the use of homogeneous solutions of purified lipids, a linear correlation has been found between the number of bisallylic methylene positions and the oxidizability of the lipids [40]. More recently, Wagner et al. [41] subjected cultured cells to oxidative stress following systematic alteration of the lipid unsaturation through supplementation of the growth medium with various PUFAs. In that study, the apparent oxidizability of the cellular lipids correlated exponentially with the number of bisallylic methylene positions in the cellular fatty acids. Different responses by the homogeneous and cellular systems to changes in PUFA content may be explained by a clustering of lipids within cell membranes that increases the apparent substrate concentration. Alternatively, Ursini et al. [42] suggested that unsaturated fatty acids are drawn into clusters of peroxidized lipids as part of a phase-compensating behavior. Such a process would “feed” the peroxyl radical–propagating reactions within the clusters of peroxidized lipid. In any event, Wagner et al. [43] found no apparent effect on the rate or extent of radical formation with fatty acid chain length, whereas Yin and Faustman [44] found with their liposomal model that both increased unsaturation and increased chain length resulted in greater phospholipid and oxymyoglobin oxidation. Location of methylene-interrupted double bonds also appears to affect the rate of oxidation as n-3 fatty acids autoxidized faster than n-6 fatty acids [45].

Table 3  Phospholipid Composition (wt %) of Subcellular Membranes in Muscle

<table>
<thead>
<tr>
<th>Phospholipid</th>
<th>Sarcolemma</th>
<th>Sarcoplasmic reticulum</th>
<th>Mitochondria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phosphatidylcholine</td>
<td>45.5</td>
<td>58.2</td>
<td>48.9</td>
</tr>
<tr>
<td>Phosphatidylethanolamine</td>
<td>22.4</td>
<td>29.4</td>
<td>39.3</td>
</tr>
<tr>
<td>Phosphatidylserine</td>
<td>17.6</td>
<td>9.4</td>
<td>9.0</td>
</tr>
<tr>
<td>Sphingomyelin</td>
<td>14.4</td>
<td>3.2</td>
<td>2.9</td>
</tr>
</tbody>
</table>

Source: Ref. 36.
Despite the appearance of oxidative products in stored muscle foods, in some studies investigators were not able to detect losses of unsaturated fatty acids [46–48]. Failure to observe measurable losses in PUFAs during frozen storage of carp may have been due to considerable fish-to-fish variation in lipid composition [49]. Even a paired-fillet technique designed to minimize the variability in triacylglycerol composition of mackerel or catfish was insufficient to detect losses in total lipids or the triacylglycerol fraction [50]. However, focusing on specific lipid fractions rather than total lipid did lead to greater sensitivity in detection of losses of unsaturated fatty acids, with losses occurring primarily in the phospholipid fraction [47,51,52].

2. Lipid Composition of Muscle

A wide degree of variation in lipid content exists among muscles from different species (Table 4) [51,53,54]. Typically, in these muscles, the quantity of phospholipids is 500 mg/100 g muscle, directly paralleling the actual amount of membrane [55]. The remainder of the lipid may therefore be considered to be primarily nonpolar triacylglycerols. However, the level of triacylglycerol does not determine the oxidative susceptibility of that sample. Rather, the relative reactivity and accessibility to catalysts and inhibitors constitute the major determinants for identification of the critical site of oxidation. Membrane lipids are distributed throughout the tissue, but the triacylglycerols or storage lipids often are not. In chicken and trout muscle, adipose cells containing the triacylglycerols were primarily found in peripheral subcutaneous fat; in red meat muscle, adipose cells were found both between and within muscle fibers; and in salmon muscle, adipose cells were mainly distributed in the myosepta and, to a lesser extent, in the connective tissue surrounding bundles of white muscle fibers [56,57]. One must keep in mind, though, that adipose cells are bounded by membrane phospholipids. Oxidation of either fraction may then spread to the other lipid site. When such spreading occurred for a peroxidizing fish microsomal fraction system, emulsified lipids added to the system were oxidized [58]. Carbon-centered lipid radicals, which have been found in the extracellular medium of oxidizing cells, may be the vehicle by which oxidation of other adjacent lipid structures occurs [59].

As shown in Table 4, the degree of polyunsaturation in the muscle varies depending on the species. To understand the importance of this component to oxi-

<table>
<thead>
<tr>
<th>Muscle/strain/variety</th>
<th>% Fat</th>
<th>% PUFA</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Channel</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aqua</td>
<td>3.7</td>
<td>24.4</td>
<td>51</td>
</tr>
<tr>
<td>LSU</td>
<td>5.4</td>
<td>22.1</td>
<td>51</td>
</tr>
<tr>
<td>Tilapia</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Red</td>
<td>1.7</td>
<td>26.1</td>
<td>53</td>
</tr>
<tr>
<td>Blue</td>
<td>1.9</td>
<td>24.5</td>
<td>53</td>
</tr>
<tr>
<td>Beef</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Longissimus dorsi</td>
<td>4.4</td>
<td>4.9</td>
<td>54</td>
</tr>
<tr>
<td>Semimembranosus</td>
<td>3.5</td>
<td>6.6</td>
<td>54</td>
</tr>
<tr>
<td>Pork</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Longissimus dorsi</td>
<td>4.5</td>
<td>5.3</td>
<td>54</td>
</tr>
<tr>
<td>Semimembranosus</td>
<td>3.3</td>
<td>9.2</td>
<td>54</td>
</tr>
<tr>
<td>Chicken</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breast</td>
<td>1.4</td>
<td>18.9</td>
<td>54</td>
</tr>
<tr>
<td>Thigh</td>
<td>6.0</td>
<td>15.5</td>
<td>54</td>
</tr>
</tbody>
</table>
dative stability, comparisons between species or strains have been conducted and related to its fatty acid composition. In some cases, such as for two strains of channel catfish, the stabilities reflected the level of polyunsaturation [51], whereas in the case of frozen raw beef, pork, or chicken, they did not [54]. Heme pigments and levels of catalase instead were suggested to be the important determinants differentiating the stabilities of those meats.

Studies involving dietary modification, on the other hand, have consistently shown that oxidative stability reflects unsaturation in the muscle tissue. When the level of unsaturation in the diet was increased, lipid oxidation occurred to a greater extent (Table 5) [60–70]. Decreased lipid oxidation has been observed with feeding diets containing increased levels of saturated, monounsaturated, or conjugated linoleic fatty acids. Feeding diets containing oxidized dietary oil may or may not affect stability. Whereas Monahan et al. [63] did not see a significant influence on oxidation in pork chops when oxidized oil was incorporated into the diet of the animal, other investigators have observed an increased oxidative lability by lipids in broilers [66,71]. In addition to altering oxidative susceptibilities, the fatty acid composition also affects the type of oxidative volatiles produced [72,73]. Using principal component analysis on data of volatile compounds in samples, Meynier et al. [73] distinguished oxidized breast muscles from turkeys fed a diet containing 6% tallow, rapeseed oil, or soya oil.

3. Susceptibility of Lipid Classes to Oxidize

In vitro studies have been valuable tools in defining the contribution of individual lipid classes (triacylglycerols, phospholipids, free fatty acids) to the oxidative stability of a food system. Through in vitro studies, free fatty acids have been shown to oxidize faster than triacylglycerols, [74] while the reactivity of membrane lipids is greater than that of emulsified triacylglycerols [75], apparently because arrangement of phospholipids in the membrane facilitates propagation. Proximity of the phospholipids to catalytic sites of oxidation (enzymatic lipid peroxidation, heme-containing compounds) also may contribute to the importance of membrane lipids

### Table 5  Studies Exploring the Influence of Diet on Muscle Lipid Composition and Oxidative Stability

<table>
<thead>
<tr>
<th>Source of muscle</th>
<th>Primary lipid source in diets</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lamb</td>
<td>Sunflower seed</td>
<td>60</td>
</tr>
<tr>
<td>Pork</td>
<td>Canola oil</td>
<td>61</td>
</tr>
<tr>
<td>Pork</td>
<td>Safflower oil or tallow</td>
<td>62</td>
</tr>
<tr>
<td>Pork</td>
<td>Oxidized corn oil</td>
<td>63</td>
</tr>
<tr>
<td>Rainbow trout</td>
<td>Fish oil or swine fat</td>
<td>64</td>
</tr>
<tr>
<td>Chicken</td>
<td>Olive oil, coconut oil, linseed oil, or partially hydrogenated soybean oil</td>
<td>65</td>
</tr>
<tr>
<td>Chicken</td>
<td>Oxidized corn oil</td>
<td>66</td>
</tr>
<tr>
<td>Chicken</td>
<td>Full-fat flax seed</td>
<td>67</td>
</tr>
<tr>
<td>Chicken</td>
<td>Fish meal</td>
<td>68</td>
</tr>
<tr>
<td>Chicken</td>
<td>Conjugated linoleic acid</td>
<td>69</td>
</tr>
<tr>
<td>Chicken</td>
<td>$\alpha$-Linolenic acid</td>
<td>70</td>
</tr>
</tbody>
</table>
in tissue oxidation along with the high degree of polyunsaturation in phospholipids [76]. Evidence that phospholipids are the major contributors to the development of warmed-over flavor (WOF) in meat from several different species of animals has been provided [77–82]. However, levels of total lipid seemed to be the major contributor to WOF in pork [77].

The method used to measure degree of oxidation may influence a study’s conclusions. For example, a greater concentration of volatiles produced by the triacylglycerol fraction often does not have the impact on flavor that a smaller concentration of volatiles produced by the phospholipid fraction would have. This response arises because solubility in the lipid and/or flavor threshold of many of the volatiles increases as the level of fat increases. Hence, Roozen et al. [83,84] demonstrated that lowering the fat content in model systems increases the chance of flavor defects by reducing the concentration of volatiles retained in the fat.

The time frame under which an investigator examines the relative contributions of lipid classes to oxidation may also be a determining factor in the reported results. At earlier stages of oxidation, the peroxide value of raw sardine fillets was attributed to preferential oxidation of phospholipids and in later stages to oxidation of triacylglycerols [85]. Igene et al. [86] also showed that triacylglycerols in model meat systems were slow to oxidize and as such did not serve as a source of oxidative products until late in storage. In contrast, Erickson [51] suggested that free fatty acids released from the triacylglycerols served as the major site of oxidation in early stages of frozen storage of channel catfish and that phospholipids were only major contributors in later stages of storage. Thus, accessibility of lipid to hydrolytic enzymes could be an important factor for determining the oxidative susceptibility of a lipid class.

4. Susceptibility to Oxidation of Membrane Lipids

Variations in oxidative susceptibility within the individual phospholipid classes can be ascribed to the nature of the polar head group (choline, ethanolamine, serine, or inositol) and the degree of fatty acid unsaturation of the individual phospholipid [44,87]. With chicken meat, Pikul and Kummerow [88] identified phosphatidylinositol as containing the highest malonaldehyde levels and largest percentage of PUFAs, followed by phosphatidylethanolamine (PE), phosphatidylserine (PS), phosphatidylcholine (PC), cardiolipin, lysophosphatidylcholine, and sphingomyelin. In model meat systems, PE consistently changed more than PC in polyunsaturation during frozen storage, reflecting the higher initial levels of unsaturated fatty acids in PE [86]. Even when fatty acid composition was held constant in a liposome model system, PE-based liposomes exhibited a greater increase in both lipid and oxymyoglobin oxidation than PC-based liposomes [44]. In contrast, saturated PS added to PC-based liposomes inhibited oxidation through modification of surface charge and subsequent trapping of iron [89,90]. Similarly, inhibition by plasmalogen phospholipids (glycerophospholipids that contain a vinyl ether moiety at the sn-1 position) has been ascribed to their binding of iron [91] and to decreased propagation via oxidation of the vinyl ether bond [92]. On the other hand, possible causes of the inhibition of membrane phospholipid oxidation upon incorporation of cholesterol include an alteration in packing of lipids in the membrane and chemical trapping of oxygen upon conversion of cholesterol into nonradical oxide derivatives [93–95].
5. Hydrolysis of Lipids and Associated Effects on Lipid Oxidation

Disintegration of lysosomal membranes in muscle tissue may occur upon mincing or storage of nonheated muscle foods. As a result, muscle lipids may be exposed to lipolytic enzymes that are released from these organelles. In support of this statement is a report that both lipase and phospholipase activities were found in frozen fish [96]. While phospholipases have been shown to be heat-inactivated more quickly than lipases [97], responses by these enzymes to frozen storage temperatures have been variable. When oyster was stored at −35°C, the activity of lipase was suppressed much more than that of phospholipase [47]. In contrast, in frozen cape hake mince, phospholipids were hydrolyzed faster than the neutral lipids above −12°C, whereas neutral lipids were hydrolyzed faster than phospholipids below −12°C [98].

As opposed to short chain free fatty acids in dairy products, long chain free fatty acids released in muscle foods do not contribute directly to rancid aromas. In general, further oxidation of these fatty acids is necessary to generate volatile products that are associated with sensory deterioration of the product. This mode of deterioration is distinct from the sensory deterioration described by Refsgaard et al. [99] for salmon. In these samples, hydrolysis of neutral lipids generated free fatty acids during frozen storage that contributed directly to an increased intensity of trained oil taste, bitterness, and metal taste by panelists.

The source of the free fatty acids determines whether lipid hydrolysis has an accelerating or inhibiting effect on subsequent rates of lipid oxidation. Free fatty acids originating from triacylglycerols accelerate oxidation [100,101], whereas free fatty acids hydrolyzed from phospholipids have been shown to inhibit oxidation [58,101,102]. In the latter case, it was suggested that free fatty acids disrupted the fatty acid alignment that facilitates free radical chain propagation in membranes [103]. Alternatively, Borowitz and Montgomery [104] concluded that the response to phospholipase may be dependent on the time of application of phospholipase. They found that when peroxidation preceded phospholipase A2 (PLA2) activity, the hydrolysis facilitated propagation of the peroxidative process. In contrast, if PLA2 was activated prior to initiation of oxidation, oxidation was inhibited. This sequence would account for the acceleration of oxidation during frozen storage when fish had been held prior to mincing [105].

While membrane lipid hydrolysis modifies the degree of lipid oxidation, the extent of hydrolytic activity may depend on the extent of membrane lipid oxidation. Oxidized fatty acids attached to phospholipids have been found to be more susceptible to hydrolysis by PLA2 than fatty acids that were not oxidized [106]. Supporting this linkage between oxidation and hydrolysis, Han and Liston [107] also found increases in both activities when ferric iron was added to fish muscle.

E. Catalysts

Catalysts of lipid oxidation in muscle foods include both enzymic and nonenzymic sources (Table 6), but by and large, the bulk of the research to date has focused on the contribution of heme and nonheme iron to promotion of lipid oxidation.

1. Transition Metal Ions

Iron heme proteins, including myoglobin and hemoglobin, are abundant in muscle tissue [108]. Relative concentrations depend on species and muscle type. Beef, lamb,
Table 6  Potential Catalysts of Lipid Oxidation in Muscle Foods

<table>
<thead>
<tr>
<th>Category</th>
<th>Catalysts</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nonenzymic</td>
<td>Transition low molecular weight metal ions</td>
</tr>
<tr>
<td></td>
<td>Metmyoglobin–H₂O₂</td>
</tr>
<tr>
<td></td>
<td>Porphyrin compounds (sensitizers for the generation of singlet oxygen)</td>
</tr>
<tr>
<td>Enzymic</td>
<td>Lipooxygenase</td>
</tr>
<tr>
<td></td>
<td>Myeloperoxidase</td>
</tr>
<tr>
<td></td>
<td>Membrane enzymic systems that reduce iron</td>
</tr>
</tbody>
</table>

and pork generally contain more myoglobin than hemoglobin, whereas chicken contains a greater amount of hemoglobin [109]. The ability of heme proteins to promote lipid peroxidation has been demonstrated by many researchers [110–112]. One mechanism of this activation involves decomposition of preformed fatty acid hydroperoxides to peroxyl radicals, which in the presence of oxygen propagate lipid peroxidation [113]. Results of experiments with inhibitors suggest that the major pathway of peroxyl radical production involves high valence state iron complexes in a reaction analogous to the classical peroxidase pathway [114]:

\[
\text{Fe}^{3+}(\text{porphyrin}) + ROOH \rightarrow \text{Fe}^{4+} = O(\text{porphyrin})^{2+} + ROH
\]

\[
\text{Fe}^{4+} = O(\text{porphyrin})^{2+} + ROOH \rightarrow \text{Fe}^{4+} = O(\text{porphyrin}) + \text{ROO}^{+} + \text{H}^{+}
\]

Heme compounds, particularly metmyoglobin and oxy-/deoxymyoglobin, also activate lipid oxidation through an intermediate species following interaction of the heme moiety with hydrogen peroxide [115–118]. An ongoing debate attempts to ascertain whether this intermediate species activates oxidation of PUFAs through ferryl oxygen [119,120], through a tyrosine peroxyl radical on the heme compound [121,122], or through radical transfer to other proteins that generate long-lived radicals [123,124]. Enhanced formation of this activated heme species has been observed when the heme compound is preincubated in the presence of the secondary lipid oxidation product 4-hydroxynonenal [125]. Decreased formation of the activated species, on the other hand, occurred in the presence of the free fatty acid linoleate [126]. In the latter case, it was proposed that metmyoglobin bound to the fatty acid anion forming a hemichrome species that could not be activated by hydrogen peroxide. However, exposure to hydrogen peroxide has also been shown to lead to release of nonheme iron from myoglobin and hemoglobin [127–129]. Alternatively, heme may donate its reducing equivalents to low molecular weight iron and copper complexes [130,131] and thereby contribute to catalysis of lipid oxidation.

In the nonheme form, iron participates in the production of the reactive oxygen species, the hydroxyl radical, via the chemical Fenton reaction: Fe²⁺ + H₂O₂ → Fe³⁺ + OH⁻ + OH⁻. This reaction is effective when Fe³⁺ can be recycled to Fe²⁺ by various reducing agents.

Levels of low molecular weight nonheme iron are initially low, being only 2.4–3.9% of total muscle iron in beef, lamb, pork, and chicken [109] and 6.7–13.9% of total iron in flounder and mackerel muscle [132]. However, in muscles that have been processed and stored, increases in the catalytic low molecular weight iron fraction have been found [132–134]. Potential sources of nonheme iron are dislodgment
of iron from the heme pocket by cooking [129,135] and release of iron from the iron storage protein, ferritin, by the reducing agents cysteine and ascorbate [136]. These increases are significant because concentrations of iron (2.2 μM) and copper (1.4 μM) that were found in the low molecular weight fractions of fish muscle were shown to catalyze lipid oxidation in fish muscle sarcoplasmic reticulum model systems [137]. In addition, modifications in dietary iron level altered the development of lipid oxidation in turkey dark muscle and in pork muscles [138–140].

Three general approaches have been taken to decipher the contribution of heme and nonheme iron to lipid oxidation of muscle foods:

1. Evaluation of the levels of heme and nonheme in the muscle food and relation of these values to the muscle’s oxidative stability during storage [54,141];
2. Evaluation of the improvement in oxidative stability upon addition of inhibitor/chelator that will cancel out the contribution of one of the components [142–145]; or
3. Evaluation of the level of oxidation induced upon addition of one of the iron sources to muscle or muscle model systems [142–148].

From these studies, the following conclusions may be drawn:

- In raw red meat and dark muscle fish, heme iron is the major catalyst.
- In both raw and cooked white flesh fish, nonheme iron is the major catalyst.

However, conflicting results make it difficult to draw a conclusion on the role of heme and nonheme iron in cooked red meat samples. Some factors that may modify the response are as follows:

1. Concentration of catalysts used in the study.
2. Distribution of catalysts in muscle system. Johns et al. [146] suggested that conflicting results from model system studies were due in part to the difficulty of evenly dispersing the catalysts in the system.
3. Concentration of reducing substances in system. Nonheme iron is more active in the reduced state, whereas heme iron is more active in the oxidized state [149].
4. pH of system. Heme catalysis is influenced less by increasing pH than nonheme iron [150]. Therefore, contribution of nonheme iron would increase with decreasing pH.
5. Amount of heat applied to the cooked system.
6. Presence of H₂O₂ in system.
7. Presence of chelators. Endogenous chelators, such as citrate, phosphate, and nucleotides, modify the reactivity of low molecular weight iron by modifying its redox potential to different extents [151].
8. Presence of one or more lipid classes. Oxidative response of chicken muscle model systems differed depending on the class of lipid(s) present [152].

2. Singlet Oxygen Generation Systems

While many different mechanisms have been proposed for the production of singlet oxygen, it is believed that the two most common mechanisms involve nonenzymatic photosensitization of natural pigments [153] or direct enzymatic production of singlet
oxygen [22]. Nonenzymatic production of singlet oxygen, in turn, involves two different pathways. The type I pathway is characterized by hydrogen atom transfer or electron transfer between an excited triplet sensitizer and a substrate, resulting in the production of free radicals or free radical ions. These free radicals may then react with triplet oxygen to produce oxidized compounds, which readily break down to form free radicals that can initiate free radical chain reactions. In the second pathway for production of nonenzymatic production of singlet oxygen (type II), the excited triplet sensitizer reacts with triplet oxygen via a triplet–triplet annihilation mechanism. Enzymatic production of singlet oxygen, on the other hand, has been shown to be a direct or indirect consequence of the action of certain microsomal oxidases, lipoxygenase, and prostaglandin synthetase.

Evidence that singlet oxygen can initiate lipid oxidation can be obtained from analysis of the oxidative products. In the reaction of singlet oxygen with unsaturated fatty acids, one end of the singlet oxygen molecule reacts with the α-olefinic carbon, while the other end abstracts the γ-allylic hydrogen. As a result of this six-membered ring transition state, both conjugated and nonconjugated hydroperoxides are formed [154], whereas free radical autoxidation of lipid produces only nonconjugated hydroperoxides.

Pigments present in muscle foods that may act as photosensitizers (because their conjugated double-bond system easily absorbs visible light energy) include hematoporphyrins and riboflavin. In model systems containing myoglobin and its derivatives, Whang and Peng [155] demonstrated through electron paramagnetic resonance spectroscopy coupled with a spin trapping technique that dissociated hematin and especially the protoporphyrin IX ring exerted a photosensitizing function. The participation of photosensitization in meat systems is supported by several storage studies: (1) for ground turkey, ground pork, and shrimp, samples exposed to light had higher levels of oxidative products than samples stored in the dark [156–158]; (2) incorporation of a UV light absorber in the packaging of pork patties prevented light-induced lipid oxidation [157]; and (3) incorporation of a singlet oxygen quencher (2,2,6,6-tetramethyl-4-piperidone) reduced the prooxidant effect of light in turkey meat [156].

3. Enzymic Initiation Systems

Several enzyme systems capable of initiating lipid oxidation have been identified in muscle foods. Among these, lipoxygenase stereoselectively absorbs a hydrogen atom from an active methylene group in 1,4-pentadiene structures of PUFA and releases a stereospecific conjugated diene hydroperoxy fatty acid product. Although it has greater recognition for its off-flavor development in vegetables and legumes, lipoxygenase has also been found in fish gill tissue [159,160], chicken muscle [161], and sardine skin [162]. However, according to Kanner et al. [163], these enzymes are not true initiators, since preformed hydroperoxides are necessary for their activation. Despite this requirement, products of lipoxygenase have been associated with “fresh flavors” of fish [160]. Their contribution to off-flavor generation in muscle foods during storage, on the other hand, remains debatable. While Grossman et al. [161] detected little loss of lipoxygenase activity in chicken muscle stored at −20°C for 12 months, German et al. [160] noted that lipoxygenases were unstable, being inactivated by 50% within 3 hours at 0°C or completely inactivated with a single freeze–thaw cycle.
Membrane systems that reduce iron constitute another enzymic system of importance in the process of lipid oxidation because they not only generate active catalysts but do so in an environment consisting of highly unsaturated membrane lipids. Membrane systems in muscle capable of generating active oxygen species in the presence of NAD(P)H and ferric iron include sarcoplasmic reticulum [164] and mitochondria [165]. Whereas the enzymic systems in beef and chicken utilize NADPH preferentially [166,167], the enzymic system in fish utilizes NADH preferentially [164]. In the latter case, both NADH-cytochrome b₅ reductase and cytochrome b₅ have been associated with reduction of low and high concentrations of ferric-histidine and low concentrations of ferric-ATP [168]. During storage, as NADH concentrations drop to levels that are maximal to the enzymic system [164], stimulation by this enzymic system would be expected to increase with time postmortem.

Myeloperoxidase is another enzyme that may be present in muscle systems postmortem and capable of initiating lipid oxidation [169]. Normally found in neutrophils of blood, myeloperoxidase may contaminate muscle tissues following slaughter and spreading of blood over the surface of the product. Even when the blood has been washed off, residual concentrations of myeloperoxidase may be sufficient to accelerate oxidation in stored foods through the following reactions:

\[
\begin{align*}
    H_2O_2 + Cl^- & \rightarrow HOCl + OH^- \\
    HOCl + O_2^- & \rightarrow O_2 + Cl^- + 'OH
\end{align*}
\]

In lipoproteins and phospholipid liposomes, hypochlorous acid (HOCl) has been found to initiate lipid oxidation [170] and may do so by interacting with organic peroxides in vivo to form reactive radicals that subsequently initiate lipid oxidation [171].

F. Antioxidants

By far the most important defense mechanism for lipid oxidation is the presence of antioxidants, which can delay or slow the rate of oxidation of autoxidizable materials. Inhibition may take two forms: a reduction in the rate at which the maximal level of oxidation is approached or a reduction in the maximal level of oxidation. This section focuses primarily on antioxidants endogenously present in muscle tissues, with only limited discussion of antioxidants applied exogenously during processing.

1. Tocopherol

The main lipid-soluble antioxidant present in muscle tissue is tocopherol. “Tocopherol” is actually used as a generic description for mono-, di-, and trimethyl tocols that contain a 6-chromanol ring structure with different numbers of methyl groups at the 5-, 7-, and 8-positions and a saturated or unsaturated 16-carbon isoprenoid side chain (Table 7). α-Tocopherol is the predominant form in muscle tissue of beef, pork, chicken, and fish, although depending on the diet composition, γ-tocopherol and α-tocotrienol may also be present to varying degrees [53,172–175]. Studies indicate that when γ-tocopherol is supplied continuously in the diet, it accumulates in the muscle but to a much smaller extent than when rats are fed similar levels of α-tocopherol [176,177]. Under these conditions, γ-tocopherol may instead tend to be accumulated in fat deposits.
Table 7  Structure of Tocopherol and Tocotrienol

<table>
<thead>
<tr>
<th>Trivial name</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>α-Tocopherol</td>
<td>5,7,8-Trimethyltocol</td>
</tr>
<tr>
<td>β-Tocopherol</td>
<td>5,8-Dimethyltocol</td>
</tr>
<tr>
<td>γ-Tocopherol</td>
<td>7,8-Dimethyltocol</td>
</tr>
<tr>
<td>δ-Tocopherol</td>
<td>8-Methyltocol</td>
</tr>
<tr>
<td>α-Tocotrienol</td>
<td>5,7,8-Trimethyltocotrienol</td>
</tr>
<tr>
<td>β-Tocotrienol</td>
<td>5,8-Dimethyltocotrienol</td>
</tr>
<tr>
<td>γ-Tocotrienol</td>
<td>7,8-Dimethyltocotrienol</td>
</tr>
<tr>
<td>δ-Tocotrienol</td>
<td>8-Methyltocotrienol</td>
</tr>
</tbody>
</table>

In general, the mechanism of antioxidant action is believed to involve competition by the antioxidant for the peroxyl radical [178]. When tocopherol acts as the antioxidant, a series of chain-breaking oxidative reactions can occur:

\[
\text{Tocopherol} \xrightarrow{\downarrow \uparrow} \text{Tocopheroxyl radical} \xrightarrow{\downarrow \uparrow} \text{Tocopherone} \xrightarrow{\downarrow} \text{Tocopherylquinone}
\]

In this mechanism, α-tocopherol donates a phenolic hydrogen atom to a lipid peroxyl radical, which forms relatively unreactive intermediates. While tocopherylquinone is a stable oxidized product, the tocopheroxyl radical and tocopherone may be reduced back to tocopherol by ascorbic acid [179] and ubiquinone [180]. Other mechanisms postulated to be involved in tocopherol's antioxidant action including scavenging of carbon-centered and hydrogen radicals [181], scavenging of singlet oxygen [182], and complexation of iron in the presence of ascorbate [183]. More recently, another mechanism was proposed for the action of tocopherol during lipid oxidation. Using egg liposomes, Dmitriev et al. [184] suggested that traditional products, such as detoxified hydroperoxides, are accumulated as well as unoxidized lipid molecules. The proposed reaction involves reduction of the peroxyl radical back to the lipid molecule, with the formation of the oxidized tocopherol and either molecular oxygen or the superoxide anion radical.

There are two approaches to identifying the contribution of tocopherol to muscle stability: monitoring of tocopherol concentrations in stored samples, and comparison of muscle stability of animals that have had dietary tocopherol supplements and animals that have not. Pfalzgraf et al. [185] incorporated both approaches into their study but found that whereas supplemented samples had improved stability compared to unsupplemented (basal) samples, α-tocopherol levels in muscle tissue did not change during refrigerated storage for either supplemented or basal samples. In contrast, losses of tocopherol have been observed in frozen pork samples [186]. Similarly, losses have been observed in frozen fish samples following a lag in early
phases of storage [51,187,188]. In these studies, accelerated degradation of tocopherol corresponded to an increased production of oxidative products. That a shift in rate of tocopherol degradation and oxidative product generation occurred within similar time periods suggested that when a critical concentration of tocopherol was reached, effective competition by tocopherol for peroxyl radicals was no longer possible and the oxidative propagation of lipids proceeded unchecked. That critical concentration varied depending on the level of phospholipid PUFAs present in the sample. Likewise, fatty acid and tocopherol compositions were considered to be the determinant factors for oxidative stability in an in vitro system [189]. Consequently, since fatty acid composition does not change in response to varying tocopherol levels [190–193], dietary supplementation with tocopherol has been found overwhelmingly to improve oxidative stability of both lipids and proteins in muscle foods as a result of deposition of the dietary tocopherol within cellular membranes [191–204]. Post-mortem supplementation of tocopherol, in contrast, has not proven as effective as dietary supplementation when levels in tissue were comparable [205], since exogenous tocopherol does not reside in the tissue membranes. However, deposition via dietary supplementation is not uniform throughout the muscles. Levels of α-tocopherol were found in beef to be highest in oxidative muscles (m. psoas major and m. glutaeus medius) and lowest in glycolytic muscles (m. longissimus thoracis and m. longissimus lumborum) [206]. In turkey, α-tocopherol levels were higher in leg muscle than in breast muscle [207], whereas in salmon they were higher in the ventral area than in the midline [208], and in herring they were lower under the skin than in other parts of the fillet [34]. The degree to which tocopherol exerts an antioxidant effect consequently depends on other compositional parameters of the muscle. For example, oxidative stabilities were improved in breast meat but not in thigh meat when chickens were supplemented with α-tocopheryl acetate [209]. The variable deposition of tocopherol homologs into muscle tissue will also factor into the changes in oxidative stability that are measured in response to dietary supplementation (α-tocopherol is preferentially deposited compared to γ- and δ-tocopherol) [210]). For example, a decreased improvement in oxidative stability was found when diets included a natural source of tocopherol (RRR-α, γ-, and δ-tocopherol) compared with a synthetic source (all-rac-α-tocopherol) [211].

2. Ascorbic Acid

Exogenous addition of ascorbic acid and its derivatives controls rancidity in a number of muscle tissues [212–217]. Within these tissues, ascorbic acid may function as an antioxidant through a variety of mechanisms: it may act as an oxygen scavenger [218]; it may scavenge free radicals generated in the aqueous phase [219]; it may maintain heme compounds in a reduced noncatalytic state [220]; and it may regenerate tocopherol [221]. At the same time, ascorbic acid can act as a prooxidant by maintaining Fe(II) in its reduced state [220]. In an investigation on ground mullet tissue, Deng et al. [222] found that ascorbic acid tended to function as a prooxidant with small quantities and an antioxidant at high concentrations, with dark muscle requiring lower concentrations of ascorbic acid for the shift to occur than light muscle. Hence, dietary supplementation with ascorbic acid would only be advantageous if the levels increase in the tissue beyond these critical concentrations. In fact, several investigators have found the absence of any enhancement in oxidative stability for broilers fed ascorbic acid-supplemented diets [202,223]. Enhancement by dietary
ascorbic acid, on the other hand, has been found when the diet also contained an elevated level of \( \alpha \)-tocopherol [223,224]. In another study, Erickson [225] demonstrated how distribution site can impact effectiveness of ascorbic acid. In that study, vacuum tumbling was used for the exogenous application of the antioxidant for intercellular distribution in channel catfish fillets, whereas an ascorbic acid bath was used to deliver the antioxidant to live channel catfish for absorption and intracellular distribution in the muscle tissue [226]. Final muscle concentrations of ascorbic acid with both treatments were twice what they were prior to treatment, but responses differed. In the case of intercellularly distributed ascorbic acid, the reducing agent protected membrane phospholipid but accelerated oxidation of triacylglycerols. In contrast, intracellularly distributed ascorbic acid did not accelerate oxidation of triacylglycerols but again protected membrane phospholipids. Considerations of compartmentation as well as the antioxidant polarity and target membrane charge must therefore be taken into account to develop more effective treatments for enhancement of tissue stability.

3. Carotenoids

The carotenoids are a group of fat-soluble pigments characterized by a linear, long chain polyene structure. Besides their roles in pigmentation and vitamin A activity, carotenoids may have a function similar to that of \( \alpha \)-tocopherol, i.e., to protect tissues from oxidative damage through scavenging of singlet oxygen and scavenging of peroxyl radicals [178]. In a manner exclusive of hydrogen abstraction, carotenoids are postulated to scavenge peroxyl radical through addition of the radical to the conjugated system such that the resulting carbon-centered radical is stabilized by resonance. When oxygen concentrations are low, a second peroxyl radical is added to the carbon-centered radical to produce a nonradical polar product. At high oxygen pressures, however, carotenoids act as prooxidants because the carbon-centered radical may add oxygen in a reversible reaction, resulting in an unstable chain-carrying peroxyl radical, which can further degrade to radicals and nonradical polar products with no net inhibition of oxidation [227]. As to performance of carotenoids in muscle tissue, the response to supplementation has varied. In postmortem supplementation, Lee and Lillard [228] observed similar levels of oxidation in cooked control and \( \beta \)-carotene-mixed hamburger patties following refrigerated storage. On the other hand, Clark et al. [229] reported that dietary canthaxanthin delayed formation of oxidative products in minced trout flesh during refrigerated storage. Similarly, Bjerking and Johnsen [230] observed a positive antioxidant effect in rainbow trout fillets. However, dietary supplementation has not always had a positive effect. Sigurgisladottir et al. [231] and Maraschiello et al. [232] did not observe any antioxidant effects by dietary carotenoids in salmon and broiler muscle tissue, respectively. Ruiz et al. [233] contended that sufficient levels of \( \alpha \)-tocopherol must be present to demonstrate an antioxidant effect by carotenoids. In cases where the proportion of carotenoid/\( \alpha \)-tocopherol is too high, the carotenoid competes with tocopherol for absorption and the levels of \( \alpha \)-tocopherol in the muscle decline. Variability in distribution of the carotenoid in the muscle tissue [208] could also account for different responses in dietary supplement studies if sample size is inadequate. As a final note, deposited carotenoids in muscle have different stabilities during storage (i.e., astaxanthin is more stable than canthaxanthin [234]), and these stabilities should be considered in supplementation of muscles to increase oxidative stability of the tissue.
4. Glutathione

While the traditional approach to understanding the role of glutathione in relation to lipid oxidation has focused on its action as a substrate for detoxification enzymes, evidence is accumulating that glutathione in and of itself acts to control lipid oxidation in several ways. In the first case, glutathione may reduce the initiator, ferryl myoglobin, back to metmyoglobin [235]. In the second case, glutathione may serve to reduce oxidized sulphydryl groups nonenzymatically [236]. Based on standard one-electron reduction potentials, Buettner [237] contends that it is thermodynamically feasible for glutathione to be oxidized to glutathione disulfide nonenzymatically, with the simultaneous reduction of a hydroxy, peroxo, or lipid radical to a hydroperoxide. Buettner’s pecking order of oxidative activity [237] is supported by the finding that in frozen minced fish, glutathione declined at a faster rate than did ascorbic acid, which in turn declined faster than \( \alpha \)-tocopherol [238]. Similarly, glutathione and ascorbate declined faster than \( \alpha \)-tocopherol and ubiquinone in both light and dark muscle of mackerel [239]. However, caution must be taken in viewing glutathione degradation uniquely as a response to inhibition because in the oxidation of glutathione to its disulfide, superoxide anions can be produced. If the superoxide anion is not removed quickly through the action of superoxide dismutase, the net effect of the reduction of harmful radicals may be minimal owing to the formation of an active oxygen species that could lead to initiation or promotion of propagation in lipid oxidation. Alternatively, in the presence of low concentrations of oxygen, thiol radicals could have a greater tendency to react with each other such that oxidized glutathione formation without the intermediate production of superoxide anion and hydrogen peroxide could occur.

The extent to which glutathione is capable of inhibiting cellular lipid oxidation in muscle foods is questionable. While Murai et al. [240] indicated that reduced glutathione was effective in limiting the adverse effects of oxidized fish oil in the diet of yellowtail fish, dietary supplementation of fingerling channel catfish did not affect fish performance, body composition, or stability of fillet samples [241].

5. Carnosine

Carnosine (\( \beta \)-alanyl-L-histidine) is an endogenously synthesized dipeptide present in beef, pork, chicken, and fish skeletal muscle at concentrations ranging from 0 to 70 mM [242–244]. Its exogenous addition to salted ground pork and beef muscle inhibited lipid oxidation [242,246], supporting model system studies demonstrating the inhibition of oxidation promoted by iron [247], hydrogen peroxide–activated hemoglobin [247], lipoygenase [247], singlet oxygen [248], peroxyl radicals [249,250], and hydroxyl radicals [251]. Inhibitory action by carnosine may be related to its ability to chelate copper ions [252], scavenge free radicals [253], or trap volatile aldehydes [254,255]. While carnosine at a 0.09% level in the diet has increased the oxidative stability of muscle [256], its high prices makes its use as a feed additive impractical. Supplementation of diets with the carnosine precursors histidine and \( \beta \)-alanine has been attempted; however, it did not prove to be an efficient method for improving the oxidative stability of pork [257].

6. Flavonoids and Phenolic Acids

Flavonoids are secondary products of plant metabolism and include more than 4,000 individual compounds divided into six subclasses: flavones, flavonones, isoflavones,
flavonols, flavanols, and anthocyanins. Phenolic acids, structurally related to flavonoids, serve as precursors of flavonoid biosynthesis. Phenolic acids include hydroxycinnamic (caffeic, p-coumaric, ferulic, and sinapic acids), hydroxycoumarin (sco-poletin), and hydroxybenzoic acids (4-hydroxybenzoic, ellagic, gallic, gentisic, protocatechuic, salicylic, and vanillic acids). The effectiveness of flavonoids and phenolic acids in retarding lipid oxidation in foods appears to be related not only to their chelating capacity but to their ability to act as free radical acceptors. However, economic and regulatory hurdles prevent the use of the purified forms of these phenolics; consequently, plant extracts are often examined for their antioxidant potential. Examples of these types of antioxidant studies are as follows: ground green tea or commercial tea extracts in a fish meat model system [258]; ground pepper in ground pork [259]; onions in cook-chill chicken [260]; dried spices or the ethanol extract of those spices in cooked minced meat patties [261]; aloe vera, fenugreek, ginseng, mustard, rosemary, sage, and tea catechins in pork patties [262]; rosemary, tea, or coffee extracts in dehydrated chicken meat [263]; galangal (a rhizome closely related to ginger) in minced beef [264]; and potato peels, fenugreek seeks, and ginger rhizomes in ground beef patties [265]. In membrane model system studies, antioxidant efficiency of flavonoids has been found to be dependent not only on their redox properties but also on their ability to interact with biomembranes [212]. Hence, similar to tocopherol, dietary supplementation (as opposed to exogenous application of these flavonoids) may prove to be more effective. Tea catechins supplemented at a level of 200 mg/kg chicken feed were equally effective in antioxidant potential as \( \alpha \)-tocopherol at the same level for up to 3 months of frozen storage [267].

7. Antioxidant Enzymes

Superoxide dismutase, catalase, and glutathione peroxidase are enzymes present in muscle tissues that may be classified as preventive antioxidants. Superoxide dismutase converts superoxide anion to hydrogen peroxide; catalase converts the hydrogen peroxide to water and oxygen; and glutathione peroxidase converts hydroperoxides to alcohols, thereby eliminating their potential decomposition by Fe\(^{2+}\). Comparison of these enzyme activities with the metabolic activities in different fish species suggests that levels of glutathione peroxidase and superoxide dismutase reflect the degree of oxidative activity in the tissue [268]. Similarly, antioxidant enzyme activity was higher in the oxidative sartorius muscle of turkey than in the glycolytic pectoralis major muscle [269]. Nutritional status of the animal prior to slaughter may also play an important role in dictating the enzyme levels in the flesh [270,271]. For example, Maraschiello et al. [272] found that glutathione peroxidase activity in chicken thighs decreased as the level of \( \alpha \)-tocopherol in the diet increased. In contrast, homeostatic compensation did not occur for glutathione peroxidase activity in turkey muscles upon vitamin E supplementation [269]. Moreover, addition of dietary selenium was considered to be very effective in maintaining glutathione peroxidase activities in chicken muscle tissue during storage [273].

Addition of antioxidant enzymes to raw and cooked meats, on the other hand, produced only low or moderate inhibition of oxidative activity [274,275]. Such limited usefulness may be due to the susceptibility of these enzymes to inactivation by active oxygen species [276,277].
G. Mathematical Modeling

The pathways and relationships involved in lipid oxidation are complex. Consequently, there is a need to have methods to quantitatively link product composition to oxidative stability. Mathematical modeling is a tool that allows the synthesis of data from one or many experiments into an integrated system from which quantitative changes in many components may be calculated. Critical to any mathematical model is the endpoint selected for shelf life. Oftentimes, this endpoint is arbitrarily selected, whereas it should be based on consumer acceptability.

Aside from studies exploring variations in moisture and oxygen concentrations [278], few studies have attempted to model lipid oxidation, and even fewer focus on lipid oxidation in muscle foods. Kurade and Baranowski [279] reported that the shelf life of frozen and minced fish meat might be predicted by measuring total iron and myoglobin levels and the time for extracted lipids to gain 1% weight. Using these three variables, deviation from the actual “shelf life” measured 7.38%. On the other hand, the estimate for frozen shelf life of fish samples obtained by Ke et al. [280] gave an average deviation of 17% when only the last variable had been used. In either case, these models are flawed as predictors of the contribution of membrane lipids to oxidative stability. From isolated model system studies, it is known that the membrane environment is a major factor in the high susceptibility of phospholipids, and extraction of the phospholipids eliminates that factor. Experimental support was provided by Ke et al. [280] who observed that the polar lipids oxidized more slowly than the neutral lipids.

Using a slightly different approach to model lipid oxidation, Tappel et al. [281] incorporated into their model some of the major chemical features associated with lipid oxidation, including peroxidizability of polyunsaturated lipids, activation of inducers and their initiation of lipid peroxidation, concurrent autoxidation, inhibition of lipid peroxidation by vitamin E, reduction of some of the hydroperoxides by glutathione peroxidase and formation of thiobarbituric acid–reactive substances (TBARS). The equations used to model the reactions were first brought into agreement with published information on these reactions by the determination of kinetic factors: activation degradation factor, inducer loss factor, antioxidant use factor, autoxidation factor, and hydroperoxide reduction factor. Subsequently, when the simulation program was applied to tissue slice and microsomal peroxidizing systems, the results of the simulation were in agreement with experimental data.

Babbs and Steiner [282] also used a computation model based on reactions involved in initiation, propagation, and termination. Their model incorporated 109 simultaneous enzymatic and free radical reactions, and rate constants were adjusted to account for the effects of phase separation of the aqueous and membrane lipid compartments. Computations from this model suggested that substantial lipid peroxidation occurred only when cellular defense mechanisms were weakened or overcome by prolonged oxidative stress. Consequently, while this model was developed to understand the contribution of free radical reactions to disease states of living organisms, useful insights may also be gleaned from it or similar models in understanding the oxidative stability of muscle food systems. In this manner, the variability that is inherent in the composition of foods may be factored into shelf life predictions, and conditions for optimal stability may be derived.
IV. EFFECT OF PROCESSING TREATMENTS ON OXIDATION

As foods are subjected to various processing treatments prior to storage, the opportunity arises to modify their pattern of oxidation. Table 8 summarizes typical responses exhibited by muscle tissue during storage following various treatments [283–351]. Each of these treatments is covered in more detail in Sec. IV.A–IV.O.

A. Rinses

Chlorine rinses on muscle food constitute one processing step that could dramatically alter the site to which lipid oxidation is directed. Chlorine rinses, based primarily on sodium hypochlorite, are commonly used to reduce microbial loads in muscle foods. Such rinses have led to incorporation of chlorine into beef, pork, chicken, and shrimp [352–355] with phospholipids incorporating more chlorine per mole of lipid than neutral lipids. Lipid chlorohydrins, formed by the reaction of HOCl with unsaturated fatty acids [356], would likely disrupt the membrane’s physical organization and reduce free radical chain oxidation of lipids [357]. This activity could be the basis for the reduced development of warmed-over flavor in cooked and stored breast patties prepared from chickens rinsed in a chlorine bath versus patties prepared from nonrinsed chickens [358]. However, the capacity to form lipid chlorohydrins in tissue samples would be diminished in samples having high concentrations of thiol and amino groups because these groups display a much greater reactivity with HOCl than unsaturated fatty acids [359,360]. Preferential scavenging of chlorine by thiols and amino groups in dark chicken meat could account for the lack of significant sensory differences in cooked and stored thigh patties prepared from nonrinsed and chlorine-rinsed chickens [358].

Owing to health concerns about trihalomethanes and other chlorination reaction products generated during interaction of organics and aqueous chlorine, efforts have been made to explore alternatives. Chlorine dioxide has been one candidate; however,

Table 8  Typical Oxidative Response by Muscle Foods to Processing Treatments

<table>
<thead>
<tr>
<th>Processing treatment</th>
<th>Typical oxidative response</th>
<th>Ref.</th>
</tr>
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<tbody>
<tr>
<td>Rinses incorporating oxidizing agents</td>
<td>Promotes</td>
<td>283,284</td>
</tr>
<tr>
<td>Washing</td>
<td>Variable</td>
<td>285–287</td>
</tr>
<tr>
<td>Skinning</td>
<td>Variable</td>
<td>34,288</td>
</tr>
<tr>
<td>Mincing</td>
<td>Promotes</td>
<td>289–297</td>
</tr>
<tr>
<td>Salting</td>
<td>Promotes</td>
<td>297–300</td>
</tr>
<tr>
<td>Curing</td>
<td>Inhibits</td>
<td>301–305</td>
</tr>
<tr>
<td>Smoking</td>
<td>Inhibits</td>
<td>306,307</td>
</tr>
<tr>
<td>Cooking</td>
<td>Promotes</td>
<td>308–320</td>
</tr>
<tr>
<td>Deep-fat frying</td>
<td>Promotes</td>
<td></td>
</tr>
<tr>
<td>High pressure</td>
<td>Promotes</td>
<td>321–323</td>
</tr>
<tr>
<td>Vacuum drying</td>
<td>Promotes</td>
<td>324,325</td>
</tr>
<tr>
<td>Irradiation</td>
<td>Promotes</td>
<td>326–333</td>
</tr>
<tr>
<td>Glazing/edible coatings</td>
<td>Inhibits</td>
<td>334–338</td>
</tr>
<tr>
<td>Freezing</td>
<td>Inhibits</td>
<td>158,339–341</td>
</tr>
<tr>
<td>Packaging</td>
<td>Inhibits</td>
<td>328,330,331,342–351</td>
</tr>
</tbody>
</table>
with this chemical, treated salmon and red grouper fillets had elevated levels of oxidative products [283]. Similarly, channel catfish fillets rinsed in ozonated water or water containing hydrogen peroxide had higher levels of oxidative products than untreated fillets [284]. Despite the elevated levels of oxidative products in both of these fish studies, the products were still considered acceptable. However, it is unclear at this time whether the increased concentration of oxidative products initially would impact the progression of lipid oxidation during storage.

B. Washing

Washing (as opposed to rinsing) is designed to remove chemical constituents of the tissue rather than microbial contaminants. Used in the process of making surimi, washing removes not only soluble proteins but fat, prooxidants, and antioxidants as well. Muscle oxidative stability ultimately depends on the relative levels of these components removed. For example, Undeland et al. [285] demonstrated that washing removed prooxidative enzymes and low molecular weight iron and copper catalysts from minced herring. Despite the removal of these components, however, washing decreased the lipid stability of the product pointing to the simultaneous removal of antioxidants. Consequently, to offset this antioxidant loss, investigators have recommended washing in antioxidant solutions [286]. Similarly, Richards et al. [287] demonstrated that an antioxidant wash improved the stability of mackerel fillets from stage 1 rigor but not stage 3 rigor. The improved response by the fresher fish was attributed to the greater removal of uncoagulated blood in those samples.

C. Skinning

Accessibility of tissues to oxygen is considered one of the most important factors contributing to oxidative instability. While filleting of fish is a common practice, it has been shown that the skin protects underlying areas from oxidation [34]. If the skin has to be removed early in the processing chain, deep skinning is an alternative to normal skinning and has been shown to improve the cold storage stability of saithe fillets [288]. Improvement is warranted since the highest rate of oxidation is observed in the under skin layer lipids [34].

D. Mincing

Mincing muscle tissue disrupts cellular integrity and exposes more of the lipids to the oxidative catalysts; it also dilutes the antioxidants and increases the exposure of the tissue to oxygen [361]. In particular, the intracellular location of the triacylglycerols could provide protection against hydrolysis in the intact muscle; but upon mincing, this protection is minimized if not eliminated. Mechanical disruption of the tissue also induces membrane lipids to form much smaller vesicles, and the increased surface area accelerates their degradation [289]. Further promotion of oxidation occurs in mechanically deboned flesh in response to release of hemoglobin and lipids from the bone marrow [290–292] and release of nonheme iron from the iron parts of the deboner [290,293]. While meat grinder wear and degree of stress applied during deboning have not led to significant variations in oxidative stability of stored sample [293,294], the head pressure used in the mechanical deboning of roaster breasts affected the chemical composition of the product and in turn its susceptibility to lipid oxidation [292].
Hot boning (prerigor excision of muscle or muscle systems from animals) and further processing of prerigor meat have economic advantages represented by reductions in refrigeration costs, space requirements, processing delays, and product turnover time. Studies generally have found that product was less susceptible to oxidation when ground prerigor than postrigor, a result that was attributed to the higher ultimate pH in prerigor ground meat [295,296]. In a case of prerigor meat that was more unstable than postrigor meat, the increased initiation reactions in the prerigor meat may have been due to higher product temperature during grinding [297].

E. Salting

Salt (sodium chloride) is added to muscle foods for a variety of reasons, such as adding flavor and inhibiting microbial growth. Nevertheless, an accelerating effect on lipid oxidation has been found with salt in a variety of meats, including beef, pork, chicken, and fish [297–300]. Although Ellis et al. [362] did not attribute the prooxidative effect of salt to the chloride ion, Osinchak et al. [363] identified chloride as the active component of salt in a liposomal model system and suggested that it may operate through release of iron from ferritin or modification in bilayer organization. Concentration of the chloride anion in the system also appears to affect the response, with low concentrations elevating and high levels inhibiting lipid oxidation. Another factor that influences the response to the chloride anion is the associated cation in the salt. Divalent cations were more stimulatory than monovalent cations at equivalent concentrations of chloride up to 0.22 M chloride [363]. Wettasinghe and Shahidi [364] concluded that mediation in response to anions by cations is through their ability to participate in ion pairing interactions with anion counterparts.

F. Application of Curing Mixtures

1. Nitrite-Based Curing Agents

Meat preservation by means of curing is typically obtained by application of mixtures containing nitrite as the key ingredient. Other ingredients in the curing mixture include sodium chloride, sugars, ascorbate, polyphosphates, and spices. Nitrite imparts multiple functional roles to cured products, inhibiting spore germination of Clostridium botulinum when added in combination with sodium chloride, producing the characteristic cured meat color, contributing to the characteristic cured meat flavor, and inhibiting the development of warmed-over flavor in cooked cured meats. Mechanisms proposed for the antioxidative activity of nitrite include formation of a strong complex with heme pigments (thereby preventing the release of nonheme iron and its subsequent catalysis of lipid oxidation), complexation of nonheme iron (which is catalytically less active than noncomplexed iron), and reaction with membrane-unsaturated lipids (which stabilizes the lipids) [301–303].

2. Nitrite-Free Curing Agents

A particular concern with the use of nitrite for the curing of meat has been the formation of N-nitrosamines, which are known carcinogens. Given the unlikelihood of finding a single compound that could perform all the functions of sodium nitrite, research directed to the elimination of the use of nitrites focused on formulation of
multicomponent alternatives [304,305]. During this search, dinitrosyl ferrohemo-
chrome was synthesized from hemin and nitric oxide and found to be capable of
imparting a characteristic cured color to meat [365]. Subsequently, this natural
cooked, cured meat pigment (CCMP) was demonstrated to accentuate the antioxidant
activity of the ingredients used for flavor preservation in nitrite-free curing compo-
sitions [305]. In a β-carotene–linoleate model system, the antioxidant properties of
CCMP were found to be concentration-dependent and were hypothesized to involve
quenching of free radicals [366].

G. Smoking

Smoking is a process that combines the effects of brining, heating, drying, and finally
application of smoke to the product. The effectiveness of smoke as an antioxidant
in processed meats is attributed to the phenols generated during thermal decompo-
sition of phenolic acids and lignin [306]. In addition, smoke flavor may mask rancid
flavor, thus requiring greater degrees of oxidation to render the product unacceptable
[307].

H. Heating/Cooking

Another processing treatment that modifies lipid oxidation is the application of heat.
Dislodgement of iron from heme compounds, disruption of cellular integrity, break-
down of preexisting hydroperoxides, and inactivation of lipases, phospholipases,
lipoxygenase, and other enzymes associated with lipid peroxidation are consequences
of heating and as a general rule lead to an acceleration in oxidation of stored pre-
cooked product. For example, heat-processed dark ground mackerel muscle oxidized
faster during refrigerated storage than its raw counterpart; however, the opposite
trend observed for the light ground muscle exemplified the exception to the rule
[308].

The response of a product to heat is dependent on the end-point temperature
and the overall amount of heat applied [285,309]. A mathematical model derived to
predict the development of warmed-over flavor in minced beef during chill storage
under various heating conditions estimated increasing levels of TBARS with increasing
end-point temperature (60–80°C) [310]. Spanier et al. [311] similarly found that
higher core temperatures (68.3°C vs. 51.7°C) in beef miniroasts caused higher levels
of TBARS. On the other hand, Smith et al. [312] determined that acceleration in
oxidation with increasing temperatures occurred only above a threshold temperature,
which in the case of chicken breast was 74°C. The existence of a threshold tempera-


ture would explain results of Mast and MacNeil [313] who pasteurized mechanically
deboned poultry at 59–60°C for up to 6 minutes and found that the treatment did
not lead to acceleration in lipid oxidation during subsequent frozen storage at −18°C.
Wang et al. [314] also found in the heating of lake herring that there existed a
breakpoint in the amount of heat applied below which inactivation of lipoxygenase-
like enzyme(s) occurred and above which factors contributing to nonenzymatic ox-
idation increased. As temperature is further increased, another breakpoint develops
in response to the generation of antioxidative Maillard reaction products. According
to Hamm [315], the Maillard reaction in meats begins at about 90°C and increases
with increased temperature and heating time. Later results support this statement:
Huang and Greene [316] reported that beef subjected to high temperatures and/or
long periods of heating developed lower TBA numbers than did samples subjected
to lower temperatures for shorter periods of time. Differences previously found in
method of cooking [317,318] therefore reflected differences in quantity of heat ap-
plicated. In other cases, differences found to be characteristic of various cooking meth-
ods reflected variability in water loss rather than differences in the extent of oxida-
tion. When TBA numbers were expressed per gram tissue, Pikul et al. [319]
calculated that microwave cooking led to lower numbers than convection oven cook-
ing. When expressed per gram of fat, however, the differences in results between
microwave and convection cooking were not significant.

Length of refrigeration prior to cooking also affects the response of muscle
tissue to cooking. Erickson [320] found that cooking of minced fish stored for 5
days generated greater amounts of either TBARS or fluorescent pigments than cooking
of product stored for 7 days. A decreased loss of \( \alpha \)-tocopherol in cooked 7-day
product compared with cooked 2- or 5-day product gave further support to the con-
clusion that lipid oxidation had been inhibited during cooking in the 7-day sample.
Change in pH, polyamines, Maillard reaction products, microbial removal of hydro-
peroxides and secondary oxidative products [367], and/or phospholipase activity
were postulated as potential factors involved in the decreased response of refrigerated
product to cooking.

I. Deep Fat Frying

A specialized form of cooking is deep fat frying, in which the product is immersed
in hot cooking oil for some period of time to totally or partially cook it. Not only
does the oil serve as the medium for heat transfer, it reacts with the protein and
carbohydrate components of the food, developing unique flavors and odors that have
definite appeal to the consumer. For breaded products, deep fat frying also sets the
batter, which binds the breading to the product surface.

A number of variables in addition to time and temperature of frying may impact
the susceptibility to oxidation of a fried muscle product. Oil quantity is one of these
variables. During frying, the product loses moisture and absorbs oil. When this ab-
sorbed oil is oxidized to any extent, it accelerates degradation of the product. Bread-
ing is another variable affecting oxidative susceptibility through attenuation of the
response to oil quality. Breading inhibits the loss of moisture and absorption of frying
oil [368], with finely ground breading material decreasing oil absorption to a greater
extent than coarsely ground material [369]. Since batter and breading are in intimate
contact with surface lipids, the catalysts, activators, and inhibitors present in the
batter and breading also have the potential to affect lipid oxidation of the fried
product.

J. High Pressure

Increasing attention has been directed in recent years to the application of high
hydrostatic pressures (up to 800 Mpa) to inactivate microorganisms [370]. Another
distinct advantage claimed for pressurization is that heat-labile compounds undergo
limited degradation compared with heat processing [371]. While high hydrostatic
pressure treatment on cod and mackerel muscle prevented hydrolysis of phospholip-
ids, a slight acceleration in lipid oxidation was noted under these conditions [321].
Similarly, oxidation of sardine lipids was accelerated by a high pressure treatment,
and its extent was related to intensity and duration of the treatment [322]. Inhibition of lipid oxidation by the addition of ethylenediaminetetraacetic acid (EDTA) to pressure-treated minced pork indicated that transition metal ions were probably released from complexes and became available to catalyze oxidation in the treated samples [323].

K. Drying

Drying has been a process applied to muscle foods for hundreds of years to extend the product’s shelf life through reduction in water and subsequent inhibition of microbial growth. Freeze drying is an extension of this process; however, the porosity and surface areas of its products are higher than those of traditional dried products and therefore oxidation of freeze-dried products occurs more readily [324]. However, reduced surface areas of products subjected to controlled low temperature vacuum dehydration minimized the increased lipid oxidation associated with freeze-dried products [325].

L. Irradiation

Irradiation is the process of subjecting materials to electromagnetic radiation or electron beams of sufficient energy levels to sever chemical bonds. While application of irradiation to muscle foods is intended to control pathogenic microorganisms, undesirable sensory changes in foods, especially development of off-flavors, may also arise, especially as the dose of radiation increases. For example, both Heath et al. [326] and Hashim et al. [327] reported that irradiation of uncooked chicken breast and thigh produced a characteristic bloody and sweet aroma that remained after the thighs were cooked but was not detectable after the breasts were cooked. These off-flavors are distinct from those associated with lipid oxidation and are believed to arise from protein oxidation. To support this statement, aldehydes, ketones, and alcohols, typical volatile classes associated with lipid oxidation, did not increase following irradiation of pork loin and pork sausage [328,329], whereas irradiation increased the production of sulfur-containing volatiles (carbon disulfide, mercaptomethane, dimethyl sulfide, methyl thioacetate, and dimethyl disulfide) in pork loin [329]. The majority of these sulfur-containing volatiles dissipate from aerobically packaged samples whereas oxidative volatiles increase [329]. Acceleration in lipid oxidation of stored irradiated samples may be circumvented by storage in vacuum packaging [328,330,331] or by antioxidant addition/supplementation to muscles [328,332,333].

M. Glazing

Glazing is a popular technique applied to fish products that will be frozen. Many types of glazes have been used, but the main glaze of commercial importance is a layer of ice, usually applied by immersing the product in water or by spraying it with water. The ice layer that is formed retards dehydration by preventing moisture from leaving the product and delays oxidation by preventing air contact with the product. However, as storage time progresses, sublimation occurs, thus decreasing effectiveness of such glazes as inhibitors. Various chemicals, including disodium acid phosphate, sodium carbonate, sodium lactate, corn syrup solids, cellulose gums, and
pectinates, have at times been added to glazes so as to reduce the brittleness or the evaporative rate of the glaze; however, success has been limited [334,369].

Edible coatings (formed directly on foods) and films (preformed, then placed on foods) can also function to prevent quality losses associated with lipid oxidation by acting as oxygen barriers. Materials displaying these barrier properties include polysaccharides (alginites, pectins, agars, carrageenans, cellulose derivatives, amylose, starches, chitin, etc.) and proteins (casein, whey proteins, wheat gluten, soy proteins, corn zein, gelatin, collagen derivatives, etc.). Studies demonstrating the effectiveness of these coatings include those conducted on frozen turkey [335], frozen king salmon [336], frozen cooked ham and bacon pieces [337], and refrigerated beef patties [338]. In some cases the coatings were no more effective than ice glazes [334].

N. Freezing

An excellent method of preserving the quality of meat and fish for long periods is by freezing the product. At temperatures below \(-10^\circ\text{C}\), both enzymatic and nonenzymatic reactions associated with lipid oxidation are decreased. In the range 0 to \(-10^\circ\text{C}\), however, decreased oxidative stabilities have been noted. When water is removed as ice in this temperature range, an accelerating effect due to increased concentration of reactants is observed and it offsets the temperature-induced deceleration. Illustrating this accelerating effect of ice crystal formation on lipid oxidation, Apgar and Hultin [339] incubated microsomal membrane fractions in the presence and absence of miscible solvents (alcohols) to prevent freezing at temperatures below \(0^\circ\text{C}\). When the ratio of the reaction rate at \(6^\circ\text{C}\) was compared to that at \(-12^\circ\text{C}\), the rate of lipid oxidation was found to decrease less in the presence of ice than in the presence of alcohols.

Unless the rate is very slow, rate of freezing has been found to have little influence on the oxidative stability of frozen products [340]. Instead, storage temperatures play a dominant role in dictating the stability of muscle foods. This is particularly true when temperature fluctuations occur during storage, since the extracellular formation of ice crystals is accelerated and cellular disruption is enhanced, thus facilitating the interaction of catalysts with lipid substrates [158]. Order of time/temperature holding treatments, on the other hand, markedly influenced development of rancidity. Lamb held at temperatures of \(-5^\circ\text{C}\) to \(-10^\circ\text{C}\) before storage at \(-35^\circ\text{C}\) developed more rancidity than lamb stored at \(-35^\circ\text{C}\) first, followed by storage at \(-5^\circ\text{C}\) or \(-10^\circ\text{C}\) [341].

O. Packaging

The stability of muscle foods during storage may be influenced by the packaging system. Vacuum packaging, which restricts the oxygen concentration, has been shown in numerous studies to extend the shelf life of muscle foods [328,330,331,342–347]. Similarly, oxidation was inhibited when an oxygen scavenger was incorporated into the packaging [348]. In contrast, application of modified atmospheres has produced variable results. Nolan et al. [349] found only minimal improvement in rancidity and warmed-over flavor scores for precooked ground meat packaged in CO\(_2\) or N\(_2\) compared with packaging in air, whereas Hwang et al. [350] found that cooked beef loin slices packaged in an 80% N\(_2\) and 20% CO\(_2\) gas mixture
responded similarly to vacuum packaging. Since residual oxygen concentrations in ground meat would be expected to be greater than intact samples, the responses from these two studies support those of model systems [372], which suggest that oxygen concentrations must be extremely low (<100 mm Hg) before significant reduction in rates of lipid oxidation can be achieved.

Packaging materials present other opportunities to inhibit rates of lipid oxidation in stored samples. Antioxidants incorporated into the packaging inhibited lipid oxidation of fish muscle during refrigerated storage [351]. Similarly, packaging materials that incorporated a UV light absorber prevented light-induced lipid oxidation in pork patties [157].

V. SUMMARY

During refrigerated and frozen storage, many muscle foods are susceptible to degradation by lipid oxidative mechanisms. This chapter has reviewed the compositional factors in muscle that have been identified as contributing to or modifying lipid oxidation reactions. Interactions abound among these factors and are responsible for the inconsistencies in shelf life evaluation of muscle foods. To fill the void, a greater emphasis should be placed on generation of mathematical models that can be used to link a muscle’s oxidative stability to its composition. Ultimately, these models should be integrated with models simulating the effects of process variables. This type of approach would facilitate shelf life prediction of muscle foods and derivation of conditions for optimal stability without the need to conduct accelerated storage tests on each batch of material being processed.

REFERENCES


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267. C. Lammi-Keefe, P. B. Swan, and P. V. J. Hegarty. Effect of level of dietary protein and total or partial starvation on catalase and superoxide dismutase activity in cardiac


I. INTRODUCTION

Oxidation of fatty acids in plant tissues is basically the same as that in animal tissues. In both systems, fatty acid oxidation occurs by at least four separate pathways: β-oxidation, α-oxidation, monoxygenation, and lipid peroxidations. β-Oxidation and α-oxidation result in the catabolism of fatty acids. Enzymatic lipid peroxidation generates important biologically active compounds that have roles in cell responses to injury. Nonenzymatic lipid peroxidation can be involved in the deterioration of tissues.

There are differences between fatty acid oxidation as it occurs in plants and in animals. For example, β-oxidation is a major source of energy in animal tissues, while it is not a major metabolic process in the majority of plant tissues, being involved only in carbon flux from lipid reserves to carbohydrate in lipid-storing nutrient tissues of seeds during germination. Complete β-oxidation happens only in mitochondria of animal cells; in plant cells, thus far, it has been demonstrated only in peroxisomes. α-Oxidation appears to be required for normal function of brain tissues of animals; presently, however, the importance of α-oxidation is not well defined in plant tissues. Monoxygenation of fatty acids is a biosynthetic process, generating hydroxyl and oxo derivatives of surface lipid polymers (cutin, suberin) in plants, whereas it is evidently a part of catabolic processes in animal tissues. In animal tissues, polyunsaturated fatty acids (PUFAs) are chiefly peroxidized by autoxidation. However, in plant tissues, lipid peroxidation has been thought to be predominantly due to enzymatic processes. The lipoxygenase (LOX) pathway is the
only enzymatic pathway of fatty acid peroxidation known to be operative in plants, whereas animals have two primary enzymatic dioxygenation pathways: one initiated by LOX and the other by cyclooxygenase.

II. β-OXIDATION PATHWAY

The β-oxidation pathway involves the oxidative degradation of a saturated acyl CoA by a recurring sequence of four reactions: oxidation linked to flavin adenine dinucleotide (FAD), hydration, oxidation linked to NAD⁺, and thiolysis by CoA to form acetyl CoA. The chain is broken between α and β carbon atoms, and the fatty acyl chain is shortened by two carbon atoms as a result of these reactions. In the strict sense, straight chain, saturated, common fatty acids (carbon chain length >3) can be catabolized completely only by continuous repetition of the β-oxidation reaction sequence.

Evidence for in vivo β-oxidation pathway in plants was first indicated by Grace [1] and by Synerholm and Zimmerman [2] in experiments analogous to the classic work of Knoop [3]. In 1956 Stumpf and Barber [4] demonstrated in vitro that lipid-mobilizing tissues of germinating oilseeds degraded fatty acids by the β-oxidation pathway known for mammalian and microbial cells. The plant cell fractionation studies showed the pathway was ascribed to mitochondria. However later studies showed that in these seeds β-oxidation activity is associated primarily with the soluble protein or extramitochondrial fraction unlike the case of the mammalian system [5,6]. In 1969 both Beevers’ group [7] and Stumpf’s group [8] established that the β-oxidation pathway was located within fragile organelles, referred to as glyoxysomes, separated from the endosperm of the castor bean (Ricinus communis L.). Subsequent studies showed clearly that all the enzymes of the β-oxidation complex, together with the fatty acid activation enzyme, were associated with glyoxysomal membrane [9,10]. Following these discoveries, it was demonstrated for lipid-mobilizing tissues of germinating seeds of various species that β-oxidation was located in glyoxysomes and was involved in conversion of oil to carbohydrate via the glyoxylate bypass [11].

The first reports presenting evidence that the ability to carry out β-oxidation is not restricted to glyoxysomes among plant peroxisomes were published between 1981 and 1983 [12,13]. Since then, investigators have demonstrated β-oxidation enzymes and/or β-oxidation activity in peroxisomes isolated from photosynthetic tissue, roots, and other plant tissues/organs that were devoid of storage lipids. Moreover, recent studies established that peroxisomes are able to metabolize physiologically relevant fatty acids of different molecular structure and to degrade completely the carbon chain of these fatty acids to acetyl CoA. Thus, current experimental evidence strongly supports the concept that the peroxisomes of higher plant cells are a compartment competent for fatty acid catabolism by the β-oxidation pathway [14].

A. Peroxisomal β-Oxidation

The β-oxidation reaction sequence in higher plant peroxisomes appears to be identical to those established for the β-oxidation reaction sequence in mammalian mitochondria and in bacteria.
1. Fatty Acid Activation

Prior to degradation by β-oxidation, fatty acids must be converted to an active intermediate, acyl CoA. In higher plant peroxisomes, fatty acids can be activated by two mechanisms.

Straight chain fatty acids are usually activated by an acyl-CoA ligase (EC 6.2.1.3). Acyl-CoA ligase is a membrane-bound enzyme and is located inside plant peroxisomes [15]. As in mammalian tissues, common addition of ATP, Mg\(^{2+}\), and CoA-SH to broken peroxisomes as the enzyme source results in the formation of acyl CoA, AMP, and pyrophosphate in a 1:1:1 stoichiometry [9,15].

The activation of straight chain fatty acids by acyl-CoA ligase is dependent on chain length [9,15,16]. Optimum activity of the enzymes has mainly been observed with C\(_{12}\), C\(_{14}\), or C\(_{16}\) saturated fatty acids or C\(_{18}\) unsaturated fatty acids. Short chain fatty acids (including propionic acid) are rather poor substrates (reaction rates <30% of the optimum activity observed). However, acyl-CoA ligases seem not to be a key enzyme in the control of β-oxidation of fatty acids in plant tissues. Ricinoleic acid (\(\alpha\)-12-hydroxy-9\(\alpha\)-octadecenoic acid), which comprises more than 80% of the fatty acids esterified in the storage triacylglycerols of castor bean endosperm, is activated in glyoxysomes from this tissue at rates somewhat lower than those obtained with palmitate (16:0) as substrate (which in this case is a rather unphysiological substrate) [9,17]. Furthermore the rate of ricinoleate activation accounts for approximately 40% of the rate required to support the calculated carbon flux from fat to sucrose in the castor bean endosperm. Acyl-CoA ligases from tissues that contain appreciable amounts of oleate (18:1) esterified in their storage triacylglycerols show, in some cases, low or no activity with 18:1 as substrate. The acyl-CoA ligases of glyoxysomes isolated from rape (Brassica napus L.) cotyledons do not activate the erucic acid predominant in the fatty acid components of the reserve lipids of the rape variety [16].

Investigation of catabolism of the branched chain amino acids leucine, isoleucine, and valine by means of a β-oxidation system demonstrates the existence of a different mechanism for activation of 2-oxo fatty acids in plant peroxisomes. In the presence of a branched chain 2-oxo acid, CoA-SH, and NAD, peroxisomes from mung bean hypocotyls catalyze the formation of CO\(_{2}\), NADH, and acyl CoA [18]. The acyl CoA contains one carbon atom less than the branched chain 2-oxo acid used as substrate. The release of CO\(_{2}\) and the formation of NADH and acyl CoA occur in a 1:1:1 stoichiometry, indicating the branched chain 2-oxo acids are activated by oxidative decarboxylation. 2-Oxo acids can be formed by fatty acids with a hydroxyl or ketone group located at an even-numbered carbon atom of a straight chain fatty acid through continuous β-oxidation. Fatty acids carrying such a functional group have been found to occur in some storage triacylglycerols of oilseeds, such as ricinoleic acid in castor bean oil. It has been demonstrated that the intermediates of ricinoleic acid oxidation, 2-oxo-octanoate, are activated by the oxidative decarboxylation mechanism.

2. The β-Oxidation Reaction Sequence

The β-oxidation reaction sequence (Fig. 1) starts by oxidation of acyl CoA to 2-\(\zeta\)-enoyl-CoA. Subsequent addition of water leads to the formation of L-3-hydroxyacyl-CoA, which is then oxidized to 3-oxo-acyl-CoA. The 3-oxo-acyl-CoA is cleaved by
Figure 1  Peroxisomal β-oxidation activation and pathway in plants: 1, acyl-CoA oxidase; 2, multifunctional protein (enoyl-CoA hydratase activity); 3, multifunctional protein (3-hydroxyacyl-CoA dehydrogenase activity); 4, 3-oxo-acyl-CoA thiolase. Asterisk indicates activation of 2-oxo acid catalyzed by oxidative decarboxylation. (Adapted from Ref. 41.)

a thiolytic step with CoA-SH to acetyl CoA and an acyl CoA containing two carbon atoms less than the parent acyl CoA. The intermediates (acyl-CoA thioesters) of this reaction sequence in higher plant peroxisomes are identical to those established for the β-oxidation reaction sequence in mammalian mitochondria and in bacteria; however, thus far except for the thiolic cleavage step, the individual steps of the β-oxidation reaction sequence are catalyzed in plant peroxisomes by quite different enzymes.

The first β-oxidation step of the plant peroxisomal β-oxidation reaction sequence leading from acyl CoA to 2E-enoyl-CoA is an oxygen-dependent reaction. Reaction products are \( \text{H}_2\text{O}_2 \) [7,19] and enoyl CoA [19]. The plant acyl-CoA oxidase is a homodimer \((M, 150 \text{ kDa})\), FAD-containing protein like the acyl-CoA oxidase of mammalian and yeast peroxisomes [20]. The purified enzyme of glyoxysomes from cucumber (Cucumis sativus L.) cotyledons preferentially acts on long chain acyl CoAs. However, linoleoyl CoA and linolenoyl CoA are not as good substrates as other long chain fatty acids [20]. The enzyme from spinach (Spinacia oleracea
L.) leaf peroxisomes exhibit higher activity toward long chain fatty acids [21,22] and low activity toward short chain acyl CoAs. The acyl-CoA oxidase of mung bean hypocotyl peroxisomes, which in other respects exhibits kinetic properties similar to those outlined above for the enzymes from other sources, shows an additional, pronounced optimum of activity with butyryl CoA as substrate [21–23]. Activity with butyryl CoA considerably higher than that obtained with palmitoyl CoA has also been reported for the acyl-CoA oxidase of peroxisomes from tubers of Jerusalem artichoke (Helianthus tuberosus L.) and pea (Pisum sativum L.) cotyledons [13,24]. The acyl-CoA oxidases of peroxisomes from the scutella of rape cotyledons and maize (Zea mays L.) show a very strong preference for C12 and/or C14 acyl-CoA with respect to activity [16]. Thus substrate specificity of acyl-CoA oxidases appears to depend on the enzyme sources.

The reactions leading from 2E-enoyl-CoA to L-3-hydroxyacyl-CoA and from L-3-hydroxyacyl-CoA to 3-oxo-acyl-CoA are catalyzed by one and the same protein [25], which exhibits both enoyl-CoA hydratase (EC 4.2.1.17) and L-3-hydroxyacyl-CoA dehydrogenase (EC 1.1.1.35) activity in plant peroxisomes. Behrends et al. [26], who studied glyoxysomes from cucumber cotyledons and peroxisomes from green leaves of Lens culinaris L., found two isoforms of the multifunctional protein (M, 74 and 76.5 kDa) and showed that these differ both in molecular structure and in kinetic properties. The ratio of 2-enoyl-CoA hydratase activity to L-3-hydroxyacyl-CoA dehydrogenase activity for the 76.5-kDa protein is three times higher (approximately 10.0) than for the 74-kDa protein. The 2-enoyl-CoA hydratase activity of glyoxysomes from cotton (Gossypium hirsutum L.) cotyledons decreases (100-fold) with increasing chain length of the enoyl-CoA (C4 to C16) [27]. The $K_m$ value for the enoyl CoAs of chain length C4 to C16 increased approximately 10-fold over the chain length range studied.

So far, there are no reports concerning kinetic properties of the 3-oxo-acyl-CoA thiolase (EC 2.3.1.16) of plant peroxisomes.

Although peroxisomal $\beta$-oxidation described above is based predominantly on the type of individual enzymatic reaction involved in $\beta$-oxidation instead of identification of the intermediates themselves or the products of the individual enzymatic reactions, the complete degradation of fatty acids by plant peroxisomes was demonstrated by Donaldson and Fang [28] using 16:0 substrate (concentrations <10 $\mu$M) and following product formation until it ceased. These investigators found that 7 nmol of NADH was formed per nanomole of palmitoyl CoA and that each nanomole of [U-14C]16:0 disappearing from the reaction mixture gave rise to 8 nmol of [14C]acetyl CoA.

Comparison of relative activities among $\beta$-oxidation enzymes suggests that thiolase is the rate-limiting enzyme of the reaction sequence. However, the regulation of $\beta$-oxidation is more complicated. During complete $\beta$-oxidation of long chain acyl CoAs by glyoxysomes from sunflower (Helianthus annus L.) cotyledons under non-steady state conditions (1–3 $\mu$M substrate), transient accumulation of intermediates was observed with [U-14C]16:0, [U-14C]18:1, and [U-14C] linoleate (18:2) catabolism, suggesting that complete $\beta$-oxidation of long chain acyl CoAs involves the consecutive operation of two $\beta$-oxidation systems that differ in chain length specificity. In all cases, the accumulation occurred at the short chain (C4) intermediate level. In contrast to complete degradation of 16:0 and 18:1, complete degradation of 18:2 required removal of acetyl CoA from the reaction mixture; alternatively, a medium
chain intermediate accumulated as an end product along with acetyl CoA. Under steady state conditions, 18:2 degradation by peroxisomes from cucumber cotyledons led to intermediate accumulation at the C₄ level as well [29]. Thus, fatty acid catabolism generally appears to slow down, at least at the C₄ intermediate level. This is basically in accordance with the known kinetic properties of acyl-CoA oxidase.

Two isoforms of both acyl-CoA oxidase (AOx) and thiolase, recently identified in the cucumber cotyledon glyoxysomes, exhibit substrate specificity toward long chain (AOx I, thiolase I) and short chain (AOx II, thiolase II) substrates, respectively [30]. When palmitoyl CoA was employed, it was found to inhibit AOx II but not AOx I. However, neither thiolase I nor thiolase II is affected by palmitoyl CoA. The metabolism of short chain substrates by the multifunctional protein exhibited 50% inhibition at palmitoyl-CoA concentrations of about 2 μM; but an inhibition of the metabolism of long chain substrates by the multifunctional protein became evident only at palmitoyl-CoA concentrations exceeding 10 μM. It is suggested that, in vivo, limited amounts of glyoxysomal NAD and CoA are available at first for the degradation of long chain (C12 or more) acyl CoAs; the corresponding β-oxidation systems permit cells to avoid accumulation of long chain acyl CoAs.

3. β-Oxidation Functions

The β-oxidation pathway in animals is a major source of energy, whereas in plants β-oxidation functions in carbon mobilization during postgerminative growth (Fig. 2). During germination, fatty acids hydrolyzed from the triglyceride stored in the spherules (oil bodies) in the fat-storing cells of the oilseeds pass into the glyoxysomes and are catabolized by the β-oxidation pathway. The resulting acetyl CoA is then converted into succinate by the glyoxylate cycle. The succinate then passes out of the glyoxysome and is converted into carbohydrates, which are required for the growth of the seedling until it becomes fully photosynthetic. In this case, the β-oxidation pathway is participating in gluconeogenesis.

Recent studies further demonstrate that β-oxidation is involved in the regulation of distribution of carbon pools in plant tissues. Expression of lauric acid (12:0)-ACP thioesterase (MCTE) in leaves of Brassica napus led to enhanced 12:0-acyl carrier protein (ACP) thioesterase activity and production of 12:0 in isolated chloroplasts, but no 12:0 accumulated in leaf glycerolipids [31]. Further study showed that 12:0-CoA oxidase activity, but not 16:0-CoA oxidase, was increased severalfold in developing seeds of B. napus expressing high levels of MCTE. In addition, isocitrate lyase levels were sevenfold higher in high-12:0 developing seeds. Control B. napus seed incubated with [14C]acetate incorporated almost all label into fatty acids, whereas MCTE-expressing seeds incorporated only 50% of the label into lipids, the remainder lying in range of water-soluble components, including sucrose and malate. These results suggest that a substantial portion of the 12:0 produced in high-12:0 B. napus is recycled to acetyl CoA and sucrose through β-oxidation and glyoxylate cycle pathways [32].

In addition, β-oxidation has been presumed to be involved in biosynthesis of jasmonic acid, a growth regulator in plant tissues.

B. Mitochondrial β-Oxidation

β-Oxidation activity in mammalian cells is located in two subcellular compartments: peroxisomes and mitochondria [33]. The mitochondrial pathway of fatty acid oxi-
The glyoxylate cycle in glyoxysome and the pathway of conversion of fatty acid to sucrose in the germinating oilseeds: 1, fatty acyl-CoA ligase; 2, β-oxidation; 3, citrate synthetase; 4, aconitase; 5, isocitrate lyase; 6, malate synthetase; 7, malate dehydrogenase; PEP, phosphoenolpyruvate; PGA, 3-phosphoglycerate; DiPGA, 1,3-diphosphoglycerate; FBP, fructose-1,6-bisphosphate. (Adapted from Ref. 11.)

dation represents the classic β-oxidation system. Mammalian peroxisomal β-oxidation attacks only very long chain fatty acids, and the sequence ends at octanoyl CoA. In the higher plant cells, however peroxisomes completely catabolize physiologically relevant fatty acids [20]. This finding has led researchers to ask whether higher plant cells possess an additional, mitochondrial β-oxidation system. The evidence appears to be controversial.

Experimental results that favor mitochondrial β-oxidation include the following:

1. Plant mitochondria contain L-carnitine [34,35] and carnitine acyltransferase activity [36,37]. Carnitine acyltransferase activity has been detected using long, medium and short chain acyl CoAs as substrate. The enzyme reacting with long chain acyl CoAs is thought to be involved in mitochondrial β-oxidation [38], being both associated with the outer face of the inner membrane of pea cotyledon mitochondria, as well as localized intramitochondrially [39].

2. Mitochondrial β-oxidation in animal cells is KCN-sensitive, since an acyl-CoA dehydrogenase tightly coupled to the respiratory chain catalyzes the
first oxidation step. KCN-sensitive, palmitoylcarnitine (or palmitoyl CoA plus L-carnitine)–dependent oxygen uptake by pea cotyledon mitochondria has repeatedly been observed in the presence of malate [39,40]. The KCN sensitivity of the reaction prevents peroxisomes from contaminating the mitochondrial fraction.

3. Peroxisomes do not oxidize palmitoylcarnitine. NADH formation in an NADH-dependent reaction in mitochondria from avocado (*Persea americana* L.) mesocarp or potato (*Solanum tuberosum* L.) tubers was greatly stimulated by palmitoylcarnitine [41], while L-carnitine was ineffective. However, Gerhardt [41] observes that palmitoylcarnitine has a disintegrating effect on mitochondrial membranes at micromolar concentrations, resulting in facilitated access of external substrates to intramitochondrial reaction sites.

4. Activities of β-oxidation enzymes are normally detected in mitochondrial fractions isolated from plant tissues on density gradients [13,19,42,43].

5. Enoyl-CoA hydratase and 3-hydroxyacyl-CoA dehydrogenase activities of the peroxisomal β-oxidation system are carried out by one multifunctional protein. In contrast, these activities belong to individual proteins in animal mitochondria. A protein exhibiting only enoyl-CoA hydratase activity and also differing from the peroxisomal multifunctional protein in other properties was recently isolated from pea cotyledon mitochondria [44]. In addition, antibodies raised against rat liver mitochondrial enoyl-CoA hydratase gave a positive signal in Western blots of total pea mitochondrial proteins; no signal is observed when blots of total pea peroxisomal proteins are probed.

The results that argue against mitochondrial β-oxidation include the following:

1. When potato tuber mitochondria were provided with palmitoyl-L-carnitine labeled at the C-1 position of the fatty acid moiety, no formation of [14C]acetyl CoA or labeled citrate was detected in the presence of malate, nor was there any loss of radioactivity from the substrate [41].

2. The first oxidation step of mitochondrial β-oxidation in animal cells is catalyzed by acyl-CoA dehydrogenase. All attempts to demonstrate acyl-CoA dehydrogenase activity in higher plant mitochondria using long, medium, and short chain acyl CoA’s as substrates have been unsuccessful [13,19,22,27].

3. When the ratio of enoyl-CoA hydratase, 3-hydroxyacyl-CoA dehydrogenase, or thiolase activity to the activity of the peroxisomal marker catalase is calculated, no statistically significant difference is noted between the ratios obtained for mitochondrial and peroxisomal fractions [12,45], indicating that β-oxidation enzyme activities in mitochondrial fractions might be attributed to contaminating peroxisomes rather than to mitochondrial constituents.

It appears that direct demonstration of acetyl-CoA formation from palmitoylcarnitine by mitochondria and/or of the first oxidation step of mitochondrial β-oxidation are the primary lines of evidence needed to establish the existence of a mitochondrial β-oxidation system in higher plant cells.
C.  β-Oxidation of Specific Fatty Acids

Plant fatty acids often possess other functional structures in their molecules, including double bonds in the Z configuration, hydroxyl or oxo groups at even-numbered carbon atoms, and methyl branching at certain positions of the carbon chain. These structures appear to form barriers to continuous passages of these fatty acids through the β-oxidation reaction sequence. To complete the degradation of these fatty acids, such barriers must be circumvented, and a modified β-oxidation pathway is required to allow this to happen.

1. Linoleic Acid

Naturally occurring unsaturated fatty acids usually have the double bond(s) in the Z configuration. This configuration appears to form a barrier to the degradation by the β-oxidation pathway up to the point of the double bond, yielding an intermediate that cannot be metabolized by enzymes of the regular β-oxidation reaction sequence. Linoleic acid (18:2) is 9Z,12Z-octadecadienoic acid (18:2,9Z,12Z) and contains Z double bonds at both even- and odd-numbered of fatty acid carbon atoms. The oxidation of 18:2 by plant peroxisomes is shown in Fig. 3.

The basic degradative reactions of 18:2 are the same as those for saturated fatty acids but with some additional enzymes needed. After three repetitive passages by the β-oxidation pathway, a 3Z-enoyl-CoA intermediate (3Z-dodecenoyl-CoA) is yielded when the position of the double bond at C-9 has been reached. The enoyl CoA formed by acyl-CoA oxidase and functioning as intermediate in the β-oxidation reaction sequence has, however, the 2E configuration. Conversion of 3Z-enoyl-CoA intermediate into the 2E-enoyl-CoA is accomplished by action of a Δ\(^{1}\),Δ\(^{3}\)-enoyl-CoA isomerase (EC 5.3.3.8).

The Δ\(^{1}\),Δ\(^{3}\)-enoyl-CoA isomerase of glyoxysomes from cucumber cotyledons has been purified and characterized. The enzyme is a homodimer (M\(_{r}\), 50 kDa) and catalyzes the reversible conversion of 3Z-enoyl-CoA to 2E-enoyl-CoA. Neither 2Z-enoyl-CoA, which is also a substrate for the enoyl-CoA hydratase activity of the multifunctional protein, 4Z-enoyl-CoA, nor 2E,4Z-dienoyl-CoA, a substrate of 2,4-dienoyl-CoA reductase, is accepted as a substrate by the isomerase. Besides 3Z-enoyl-CoA, 3E-enoyl-CoA is also converted to 2E-enoyl-CoA by the isomerase. Comparison of the activity of the enzyme toward 3Z- and 3E-hexenoyl-CoA indicates that the activity with the Z isomer is approximately 30 times higher. Increase of the chain length of 3E-enoyl-CoA from C\(_{6}\) to C\(_{12}\) increased the relative activity of the enzyme [46].

Further degradation of 2E,6Z-dodecadienoic acid yields a 2E,4Z-dienoyl-CoA, which is further catabolized by either 2,4-dienoyl-CoA reductase pathway or dehydratase pathway.

2,4-Dienoyl-CoA reductase (EC 1.3.1.34) reduces 2E,4Z- as well as 2E,4E-dienoyl-CoA to 3E-enoyl-CoA in an NADPH-dependent reaction. To return to the β-oxidation pathway, the 3E-enoyl-CoA formed by the 2,4-dienoyl-CoA reductase must subsequently be isomerized to 2E-enoyl-CoA by the Δ\(^{1}\),Δ\(^{3}\)-enoyl-CoA isomerase (see above). With 2E,4E-decadienoyl-CoA serving as substrate, 2,4-dienoyl-CoA reductase activity has been demonstrated in glyoxysomes from cucumber cotyledons [26] and pineapple (Ananas comosus Merr.) fruit tissue [47].

In the dehydratase pathway, 2E,4Z-dienoyl-CoA is first degraded by β-oxidation to produce a 2Z-enoyl-CoA intermediate (2Z-octenoic acid) when the position...
of the C-12 double bond has been reached. Since the 2-enoyl-CoA hydratase activity of the multifunctional protein does not show specificity for the geometrical configuration of the double bond at position 2, 2Z-enoyl-CoA intermediate is hydrated by the multifunctional protein (as is the hydration of the 2E-enoyl-CoA intermediate formed by the activity of the acyl-CoA oxidase). However, hydration of 2Z-enoyl-CoA by the enoyl-CoA hydratase reaction results in the formation of D-3-hydroxyacyl-CoA, which is not a substrate for the subsequent 3-hydroxyacyl-CoA dehydrogenase reaction. The D-3-hydroxyacyl-CoA is epimerized to its L isomer by two steps of reactions, involving the dehydration of D-3-hydroxyacyl-CoA to 2E-enoyl-CoA by a D-3-hydroxyacyl-CoA dehydratase and subsequent hydration of the 2E-enoyl-CoA to L-3-hydroxyacyl-CoA by the enoyl-CoA hydratase activity of the multifunctional protein.

In the early days, this pathway was thought to be catalyzed by a 3-hydroxyacyl-CoA epimerase (EC 5.1.2.3). Using D-3-hydroxydecanoyl-CoA as substrate, 3-hydroxyacyl-CoA epimerase activity was demonstrated in glyoxysomes of cucumber.
cotyledons and partially purified. The activity has been attributed to three distinct proteins. Two of them were the isoforms of the multifunctional protein. The third monofunctional protein carried the predominant proportion of the overall epimerase activity [29]. However, Engeland and Kindl [48] also purified two homodimeric isoforms of the novel D-3-hydroxyacyl-CoA dehydratase from cucumber cotyledons. They exhibited kinetic and molecular properties very similar to the earlier described monofunctional “epimerase” protein. Both D-3-hydroxyacyl-CoA dehydratases reversibly catalyzed the conversion of D-3-hydroxydecanoyl-CoA to 2E-decenoyl-CoA, as demonstrated by product identification, and were inactive toward L-3-hydroxydecanoyl-CoA or 2Z-decenoyl-CoA.

The following evidence indicates that the D-3-hydroxyacyl-CoA dehydratase pathway may be favored over the 2,4-dienoyl-CoA reductase pathway during 18:2 catabolism in higher plant peroxisomes.

1. Complete degradation of 18:2 as well as the rate of 18:2 catabolism were unaffected by NADPH (or NADH), which is required for participation of the 2,4-dienoyl-CoA reductase in 18:2 degradation [41].
2. The activity of the D-3-hydroxyacyl-CoA dehydratase in glyoxysomes from cucumber cotyledons is 100 times higher than the 2,4-dienoyl-CoA reductase activity.
3. The 2,4-dienoyl-CoA reductase activity amounts to only one-tenth of the activity of thiolase [48], which evidently is the rate-limiting enzyme of the β-oxidation reaction sequence in vitro.
4. The low, 2,4-dienoyl-CoA reductase activity, which should lead to intermediate accumulation at or above the C10 level, did not elicit this result [47].

Complete degradation of 18:2 by higher plant peroxisomes has been demonstrated. Other than [14C]acetyl CoA, no other products finally accumulated when peroxisomes metabolized [U-14C]18:2. The amount of [14C]acetyl CoA formed after nearly complete consumption of the 18:2 corresponds to that calculated for complete degradation of the 18:2 consumed. Under steady state conditions of [U-14C]18:2 degradation, accumulation was not observed at the C12, C10, or C8 intermediate level, where the barriers to continuous passages through the β-oxidation reaction sequence must be surmounted [26].

Based on the modified β-oxidation pathway of linoleoyl CoA, the catabolism of 18:1, linolenic acid (18:3), as well as other unsaturated, straight chain fatty acids can easily be deduced.

2. Ricinoleic Acid

An in-chain hydroxyl (or oxo) group located at an even-numbered carbon atom of a straight chain fatty acid also forms a barrier, 2-hydroxyacyl-CoA intermediate, to continuous degradation by the β-oxidation pathway. A different modified β-oxidation pathway, which allows circumvention of this barrier, is demonstrated during the complete catabolization of ricinoleate in plant peroxisomes (Fig. 4).

Following activation of ricinoleic acid by acyl-CoA ligase, ricinoleoyl CoA is degraded by β-oxidation and modified β-oxidation (see above). At the C3 intermediate level, the hydroxyl group of D-2-hydroxy-octanyl-CoA prevents repetitive passages through the β-oxidation reaction sequence. To surmount this barrier, the 2-
hydroxyacyl-CoA is first hydrolyzed by a hydrolase to yield the 2-hydroxy acid. The 2-hydroxy acid is then oxidized by a 2-hydroxy acid oxidase to form 2-oxo-octanoic acid, which is oxidatively decarboxylated to produce acyl CoA containing one carbon atom less than the 2-oxo acid. The oxidative decarboxylation allows return to the acyl-CoA track for further degradation of the parent fatty acid by β-oxidation, without the requirement of ATP for the renewed activation process [49].

The alternative reaction sequence leading from D-4-hydroxydecanoyl-CoA via 4-oxo-decanoyl-CoA to 2-oxo-octanoate has also been proposed to account for the presence of 4-oxo-decanoate, which was detected upon ricinoleate catabolism [50]. This in-chain oxidation of hydroxyl groups to oxo groups [51,52] is known as the biosynthesis of wax components (a process not located in peroxisomes).
Catabolism of heptanoyl CoA as well as ricinoleoyl CoA to propionyl CoA and acetyl CoA has been demonstrated. Following NADH formation during degradation of 1 nmol of ricinoleate, a stoichiometry of 9:1 is observed when the NADH formation has ceased. This stoichiometry corresponds to that expected for complete degradation of ricinoleate to acetyl CoA [17].

3. Propionate

Propionyl CoA can be generated at certain metabolic processes such as the catabolism of ricinoleic acid, branched chain 2-oxo acids, and odd-numbered, straight chain fatty acids by the β-oxidation pathway. In mammalian mitochondria, propionyl CoA is carboxylated to methylmalonyl CoA in a biotin-dependent reaction, and the methylmalonyl CoA is subsequently rearranged to succinyl CoA in a coenzyme B12-dependent mutase reaction. However, the propionate metabolism pathway in plants obviously differs from that in animals. Employing intact tissues from different plant species as well as cell-free systems, Stumpf and coworkers found that 3-hydroxypropionate was an intermediate of propionate catabolism; specifically, the $^{14}$CO$_2$ was released in decreasing rates from propionic-1-$^{14}$C, propionic-3-$^{14}$C, and propionic-2-$^{14}$C [53]. No Krebs cycle acids could be isolated when propionic-1-$^{14}$C was fed, whereas these acids were readily isolated when propionic-2-$^{14}$CO$_2$ and propionic-3-$^{14}$C were fed. The C-2 and C-3 of propionate became C-2 and C-1, respectively, of acetyl CoA, and CO$_2$ was not required for the degradation. On the basis of these results, the pathway of propionate catabolism called “modified β-oxidation” (Fig. 5) was proposed. This pathway avoids both a biotin-dependent carboxylation and a coenzyme B12-dependent mutase reaction and is similar to oxidation of propionate in bacteria.

Recently, catabolism of propionyl CoA to acetyl CoA via modified β-oxidation pathway was further confirmed in work that used peroxisomes from mung bean hypocotyls as the enzyme source. Gerbling and Gerhardt [49,54] used high-performance liquid chromatography (HPLC) to identify the acyl-CoA thioester intermediates, the intermediates not esterified to CoA-SH, and the end product acetyl CoA. Oxidation of propionyl CoA to acryl-CoA resulted in concomitant H$_2$O$_2$ formation, indicating that the reaction was catalyzed by acyl-CoA oxidase. In the absence of NAD$^+$, 3-hydroxypropionate accumulated as an end product. Malonic semialdehyde accumulated when CoA-SH was omitted from the reaction mixture. Catabolism of propionate via 3-hydroxypropionate to acetate was also reported in the lima bean (*Phaseolus limensis*) by Halarnkar et al. [55].

4. Branched Fatty Acids

Figure 6 shows modified β-oxidation of methyl branched fatty acids generated by the catabolism of branched chain amino acids. The β-oxidation of long chain fatty acids with methyl groups at odd- or even-numbered carbons is discussed in Sec. III.

Catabolism of leucine, isoleucine, and valine is initiated by an aminotransferase reaction yielding the branched chain 2-oxo acids, 2-oxo-4-methylpentanoic acid (2-oxo-isocaproic acid), 2-oxo-3-methylpentanoic acid (2-oxo-3-methyl-valeric acid), and 2-oxo-3-methylbutanoic acid (2-oxo-isovaleric acid), respectively. The branched 2-oxo acids are activated by the oxidative decarboxylation and then catabolized to propionyl CoA and subsequently to acetyl CoA in peroxisomes from mung bean hypocotyls. Pathways of the catabolism of the branched chain 2-oxo acids are pro-
posed based on both identification of intermediates and the results of kinetic experiments. Some individual reactions have been directly assayed. These include the oxidative decarboxylation and the subsequent reaction leading from acyl CoA to 2-enoyl-CoA. Oxidation of the branched chain acyl CoAs to the corresponding 2-enoyl-CoA resulted in concomitant H₂O₂ formation, indicating that the reaction is catalyzed by acyl-CoA oxidase [18]. It is unknown at present whether steps of the β-oxidation reaction sequence involved in the catabolism of branched chain acyl-CoAs are catalyzed by proteins identical to those acting on straight chain acyl CoAs.

III. α-OXIDATION

α-Oxidation is defined as a series of reactions by which a free fatty acid having a chain length of 13 to 18 carbons is oxidatively degraded, with the simultaneous release of CO₂ from the carboxyl carbon and the formation of a free fatty acid containing one less carbon atom (Fig. 7).
The first reaction in \( \alpha \)-oxidation is the oxidation of free fatty acids by an FAD-flavoprotein enzyme to give \( \alpha \)-hydroperoxyl fatty acids. The initial attacking species is probably the enzymatic FAD in its free radical semiquinone form (XH\(^{-}\)). This substance stereospecifically removes H\(_{\beta}\) (\( \beta \)-hydrogen atom) from the \( \alpha \)-carbon of a free fatty acid to form a fatty acid free radical and reduced flavoprotein. The reduced flavoprotein (XH\(_{2}\)) then reacts with O\(_{2}\) to form the adduct XH\(_{2}\)O\(_{2}\). The latter reacts with the fatty acid free radical resulting from the initial attack and generates a 2-\( \alpha \)-hydroperoxy fatty acid. The fatty acid hydroperoxides may then undergo anhydro-decarboxylation to yield a fatty aldehyde, as a continuation of \( \alpha \)-oxidation; or it may be reduced to form a 2-\( \alpha \)-hydroxy fatty acid as an end product. The fatty aldehyde, which is one carbon shorter than the original fatty acid, is then oxidized by an NAD-specific aldehyde dehydrogenase (EC 1.1.1.35) to its corresponding fatty acid. This completes one turn of the \( \alpha \)-oxidation spiral. Further turns successively reduce it by one carbon atom at a time [56].
The α-oxidation system was first observed in crude extracts of germinated peanut (*Arachis hypogaea* L.) cotyledons by Newcomb and Stumpf. With this microsomal system, studies from Stumpf’s laboratory [53] found several interesting results:

1. No cofactors were required for $^{14}\text{CO}_2$ production from 1-$^{14}\text{C}16:0$, but NAD$^+$ was specifically required for $^{14}\text{CO}_2$ production from internally labeled 16:0 (2-$^{14}\text{C}16:0$).
2. Synthetic 16:0-CoA was inactive.
3. The tricarboxylic acid cycle was not responsible for the CO$_2$ formation from 16:0 oxidation.
4. The system required an H$_2$O$_2$/H$^*$ generating system and a NAD$^+$-linked long chain fatty aldehyde dehydrogenase.
5. The long chain fatty aldehydes were only intermediates.
6. Addition of long chain fatty aldehydes to peanut extract resulted in the formation of the corresponding free fatty acids.

Thus, Martin and Stumpf [57] proposed the α-oxidation pathway, consisting of two separate reactions: (1) a peroxidative decarboxylation of the acid to yield CO$_2$ and an aldehyde and (2) an NAD-specific dehydrogenation of the aldehyde to the corresponding acid having one less carbon atom than the original acid.

Hitchcock and James [58] observed that 16:0 was also converted to pentadecanoate by pea leaf extracts with the accumulation of α-hydroxy fatty acids, mainly the D configuration. The leaf system required O$_2$ rather than H$_2$O$_2$. Addition of a mixture of l-D-α-hydroxy fatty acids (16:0) increased CO$_2$ release with formation of C$_{n-1}$ fatty acids. The plant leaves had a system that appeared to differ from the peanut system, and an amplified version was proposed by Hitchcock and James [59].

Apparent, differences between the two earlier proposals were reconciled by Shine and Stumpf [56], who used the same experimental system to compare partially purified preparations from pea leaves and peanut cotyledons. This investigation dem-
onstrated that α-oxidations require both oxygen and a source of reducing power. Hydroxy fatty acid was not an intermediate of α-oxidation spiral (Fig. 7).

Since the work by Shine and Stumpf, the mechanism of α-oxidation in plants has not been specifically studied, although Galliard and Matthew [60] noted that α-oxidation in cucumber fruits was not stimulated by flavoproteins and required a metal enzyme.

The α-oxidation complex is generally thought to be membrane-bound, probably to the endoplasmic reticulum [56], and has not been purified. Specificity studies [57,60,61] show that myristic and 16:0 and unsaturated C18 fatty acids are the most effective substrates (C12 << C14 > C16 >> C18; 14:0 = 18:1 = 18:2 = 18:3). Neither methyl 16:0 nor Δ216:1 is an active substrate, and 2-L-hydroxy 16:0 inhibits α-oxidation.

The physiological significance of the α-oxidation pathway remains to be carefully evaluated. Although the α-oxidation pathway produces NADH, the present consensus is that α-oxidation is not linked to ATP production.

The α-oxidation pathway may be a very effective mechanism for bypassing blocking groups in a hydrocarbon chain of a fatty acid by shifting the β-oxidation reading frame by a C1 unit, thereby allowing effective β-oxidation to occur. The involvement of α- and β-oxidation in the metabolism of phytic acid was postulated by Steinberg and colleagues [62] based on the studies on animal tissues. In patients with Refsum’s disease, an inheritable condition affecting the nervous system, large amounts of phytanic acid accumulate in tissue with a marked decrease in α-oxidation capacity, while in tissues of normal individuals, phytic acid is rapidly metabolized, although β-oxidation of pristanic acid is normal in both types of tissue. Phytic acid is 3,7,11,15-tetramethylhexadecanoic acid. The 3-methyl group completely blocks normal β-oxidation. If the reading frame is moved by one carbon, α-oxidation converts phytic acid to 2,6,10,14-tetramethylpentadecanoic acid or pristanic acid. The α-methyl group now is readily bypassed by the β-oxidation enzyme, and propionyl CoA is eliminated by the first turn of the spiral instead of acetyl CoA. Moreover, the two remaining methyl branches of the molecule also reside on α-carbon atoms as the β-oxidation spiral continues and they, too, are bypassed, with production of propionyl CoA.

2-D-Hydroxy fatty acids, a product of an offshoot of α-oxidation, are the major acyl components of cerebrosides in higher plants. α-Oxidation of even chain length acids could be the source of the odd-numbered fatty acids in some plant lipids. When an odd chain fatty acid undergoes β-oxidation, the terminal product will be propionyl CoA, which can then enter the modified β-oxidation pathway. An intermediate of the latter pathway is malonic semialdehyde, a precursor of β-alanine [63]. Thus, α-oxidation is indirectly involved in synthesis of a component of CoA and ACP.

A common α-oxidation process exists in seeds and in leaves. Laties et al. [64] found that the respiration induced by cutting potato tuber tissue was mainly due to α-oxidation of fatty acids, with carbohydrate-based respiration becoming important only at later stages of the aging process [65]. In young leaf tissue, added fatty acids were more rapidly oxidized by α-oxidation than by β-oxidation [66]. Long chain aldehydes are components of volatile products formed by plants, particularly in response to damage. The profile of the long chain aldehydes obtained from cucumber fruits was explained by the specificity of the α-oxidation system in this tissue [60].
IV. MONOXYGENASE PATHWAYS

A. \( \omega \)-Oxidation

Fatty acids with oxygen functional group (alcoholic, carbonyl or carboxyl) at the methyl terminal end (\( \omega \)-end) are formed by \( \omega \)-oxidation and frequently occur as components of cutin and suberin. The initial oxidation step that brings about the \( \omega \)-hydroxylation of a fatty acid is catalyzed by a monoxygenase. The \( \omega \)-hydroxy fatty acid is then catalyzed by NADP-specific dehydrogenases to form oxo-fatty acid and dicarboxyl acid (Fig. 8).

The hydroxylative reaction in the \( \omega \)-oxidation pathway was first demonstrated in plant tissues by Soliday and Kolattukudy [67]. In the presence of NADPH and \( \mathrm{O}_2 \), the endoplasmic reticulum fraction from embryonic shoots of \textit{Vicia faba} converted 16:0 to \( \omega \)-hydroxy 16:0. Microsomal preparations from the root tips and the excised epidermis of expanding leaves of \textit{V. faba} also catalyzed a similar hydroxylation reaction. Neither acyl CoA nor fatty acid ester was used in the oxidation. In addition to 16:0, other fatty acids especially 18:1 were a good substrate for the reaction. The \( \omega \)-hydroxylation catalyzed by the microsomal preparation was inhibited by classic mixed-function oxidase inhibitors such as metal ion chelators, NaN3, and thiol-directed reagents. The involvement of a cytochrome (cyt) P450-type protein in the \( \omega \)-hydroxylation was suggested by the inhibition of this reaction by CO. The degree of sensitivity to CO and lack of photoreversibility by light, however, implied

\[
\begin{align*}
\text{Fatty acid} & \quad \overset{\mathrm{O}_2 + \text{NADPH} + \mathrm{H}^+}{\longrightarrow} \quad \text{Cyt P-450 monoxygenase} \\
\text{OH} & \quad \text{CH}_2=\text{CH}_2(\text{CH}_2)_n\text{-COOH} \\
\text{Cyt P-450 monoxygenase} & \quad \overset{\mathrm{OH}}{\longrightarrow} \quad \text{CH}_2=\text{CH}_2(\text{CH}_2)_n\text{-COOH} \\
\text{NADP}^+ & \quad \overset{\omega \text{-Hydroxy fatty acid dehydrogenase}}{\longrightarrow} \quad \text{CH}-\text{CH}_2(\text{CH}_2)_n\text{-COOH} \\
\text{NADPH} + \mathrm{H}^+ & \quad \overset{\omega \text{-Oxo fatty acid dehydrogenase}}{\longrightarrow} \quad \text{HOOC-CH}_2(\text{CH}_2)_n\text{-COOH} \\
\text{NADP}^+ & \quad \text{Dicarboxylic fatty acid}
\end{align*}
\]

\textbf{Figure 8} The \( \omega \)-oxidation pathway in plant tissues.
that the member of the enzyme family involved was not the same one participating
in in-chain oxidation or midchain hydroxylation.

The presence of 16-oxo-9- or 16-oxo-10-hydroxypalmitic acid in cutin and
dicarboxylic acids in suberin has suggested that the ω-hydroxyl group is further
oxidized in many plant tissues. Conversion of exogenous labeled 10,16-dihydroxy-
palmitic acid to the corresponding 16-oxo acid was demonstrated with slices of
embryonic shoots of V. faba [68]. Experiments to determine how dicarboxylic fatty
acids are synthesized were carried out with cell-free extracts of epidermis of V. faba
leaves. Kolattukudy et al. [69] showed that the ω-hydroxy C_{16} acid was converted
into both 1,16-dioic and 16-oxo-C_{16} acids, while feeding the latter resulted only in
synthesis of the 1,16-dioic C_{16} acid. The dehydrogenase activity, located mainly in
the 100,000g supernatant, showed an optimum pH near 8 and a requirement for
NADP^+. Modification of substrate by esterification of the carboxyl group, replace-
ment of the carboxyl group by a methyl group, or introduction of another hydroxyl
group at C-10, rendered it a poor substrate. Thiol-directed reagents strongly inhibited
oxidation of the ω-hydroxy group. Agrawal and Kolattukudy [70] demonstrated that
two different dehydrogenases in extracts of suberizing potato slices were involved
in the oxidation of ω-hydroxy acid. One (ω-hydroxy acid dehydrogenase) converted
the C_{16} ω-hydroxy substrate to the ω-oxo derivative, and the second (ω-oxo acid
dehydrogenase) converted the latter to the dioic fatty acid. Only the ω-hydroxy acid
dehydrogenase was induced by wounding (16-fold). Its purification identified a dimer
of 31-kDa subunits. A detailed characterization revealed that this enzyme was quite
similar to other dehydrogenases [51,72]. With alkanals as model substrates, the Mi-
chaelis constant $K_m$ decreased drastically from 7000 µM to 90 µM as the chain length
of the substrate increased from C_{3} to C_{8}; a further increase in the chain length from
C_{8} to C_{20} resulted in only a small further decrease in $K_m$. Aliphatic chains longer
than C_{20} show extremely low rates.

B. In-Chain Oxidation/Midchain Hydroxylation

10,16-Dihydroxy C_{16} acid is a major cutin monomer. The hydroxylation of ω-hy-
droxypalmitic acid at C-10 was first demonstrated by Walton and Kolattukudy [71]
in a cell-free preparation from excised epidermis of expanding V. faba leaves. The
C-10 hydroxylase appeared to be located in the endoplasmic reticulum [72]. Further
purification of the crude microsomal preparation showed that the enzyme catalyzed
synthesis of both 9- and 10-isomers. This midchain hydroxylation required O_2 and
NADPH and was inhibited by NaN_3, metal ion chelators, as well as by thiol-directed
reagents. The inhibition of midchain hydroxylation caused by CO was photore-
versible, indicating a typical cyt P450 enzyme. Loss of only one hydrogen during
the hydroxylation suggested that an insertion mechanism was involved.

\[
\text{CH}_2\text{OH} \rightleftharpoons \text{(CH}_2)\text{H} \rightarrow \text{CH}_2\text{OH} \rightleftharpoons \text{(CH}_2)\text{H} \rightarrow \text{COOH} + \text{O}_2 + \text{NADPH} + \text{H}^+ \\
\text{ cyt P450 monooxygenase}
\]

\[
\text{(midchain hydroxylase)} \rightarrow \text{CH}_2\text{OH} \rightleftharpoons \text{(CH}_2)\text{H} \rightarrow \text{CHOH} \rightarrow \text{(CH}_2)\text{H} \rightarrow \text{COOH} + \text{H}_2\text{O} + \text{NADP}^+
\]

A microsomal preparation from aged Jerusalem artichoke slices catalyzed hy-
droxylation of C_{12} acid at the C-8, C-9, and C-10 positions with NADPH and O_2 as
the required cofactors [73]. This activity involving cyt P450 was absent in fresh tissue and was induced by aging the disks for 3 days. The biological significance of this observation and the relevance of choosing C₁₂ acid as the substrate are not clear.

A different enzyme appears to be responsible for the formation of hydroxy groups in ricinoleic acid from 18:1 in the developing endosperm of the castor bean. In elegant double-labeling studies, Morris [74] established that the hydrogen is directly replaced by a hydroxy group rather than via an unsaturated, keto, or epoxy intermediate. In this work, developing endosperm slices were incubated with a mixture of [1-¹⁴C]oleic acid and erythro-12,13-ditritio-oleic acid. The H/¹⁴C ratio of ricinoleate synthesized was 0.75. This result can be obtained only by a hydroxyl substitution mechanism. For a keto intermediate, it should be 50% and 50%; for an unsaturated intermediate, 50% and 25%; and for an epoxy intermediate, 50% and 25%. Hydroxylation used oleoyl CoA and required NAD(P)H and O₂ [75]. Investigation of the substrate specificity of the hydroxylase showed that chain length and double-bond position with respect to both ends were important and caused reduced activity relative to 18:1 [78]. However, the position at which the hydroxyl was introduced was determined by the position of the double bond, always being three carbons distal toward the ω end. The substrate for hydroxylation in vivo is most likely 18:1 esterified in the sn-2 position of phosphatidylcholine, from which ricinoleate was released as the free acid before activation and incorporation into triacylglycerol [77]. Antibodies raised against purified plant cytochrome b₅ inhibited the hydroxylation reaction, pointing to cytochrome b₅ as the electron donor to the hydroxylase. Carbon monoxide did not inhibit hydroxylation, and the characteristics of the hydroxylase were generally similar to those of the microsomal fatty acid desaturases. Van de Loo et al. [78] capitalized on the assumed homology to desaturases to clone the 12-hydroxylase gene from developing castor endosperm.

V. LIPID/FATTY ACID PEROXIDATION

Peroxidation is a chemical reaction that results in the generation of peroxides and their degradation products. One of the most typical properties of the unsaturated fatty acids, particularly the polyenoic acids, is their susceptibility to peroxidation. In animal tissues, PUFAAs are chiefly peroxidized by autoxidation. In plant tissues, however, lipid peroxidation has been thought to be predominantly a result of enzymatic processes, such as those of the oxylipin/LOX pathway. Lipid peroxidation produces a number of fatty acid derivatives and directly impacts various aspects of plant food product qualities.

A. Oxylipin/Lipoxygenase Pathway

“Oxylipins” is the generic name for a family of oxygenated compounds formed from fatty acids by enzymatic reaction(s) involving at least one step of dioxygen-dependent oxidation [79]. In mammalian tissues, the C₂₀ PUFA, arachidonic acid (20:4), serves as the most important precursor of oxylipins, while plant oxylipins are mainly formed from the C₁₈ PUFA 18:2 and 18:3. Oxylipins include fatty acid hydroperoxides, hydroxy fatty acids, epoxy fatty acids, keto fatty acids, volatile aldehydes, and cyclic compounds such as jasmonic acid. Although a number of interesting hydroperoxide metabolites have been defined in detail with respect to structure, an explanation of their physiological function appears to be less than complete.
Plant oxylipin pathway, also named as LOX pathway, starts with the LOX-catalyzed peroxygenation of 18:2 or 18:3 to both enantiomeric and stereospecific hydroperoxides of fatty acids. The hydroperoxides further serve as substrates for several pathways: the hydroperoxide lyase (HPLS) pathway, the hydroperoxide dehydratase (HPDS) pathway, the hydroperoxide isomerase pathway (HPIS), and the hydroperoxide-dependent peroxygenase (HPPR)/epoxygenase (HPEP) pathway. Conversion of the hydroperoxides by these pathways results in the formation of various oxylipins, including regulatory molecules and some other substances of importance in plant defense reactions.

Discovery of the plant oxylipin pathway can be traced to the early 1930s [80]. Food processing led to the first references to LOX concerning the bleaching of wheat (Triticum aestivum L.) flour and dough following the addition of small amounts of soybean [Glycine soya (L.) Merr.] flour. Since then, the impact of oxylipin pathway on foods has been of great interest to food scientists [81–84]. We now know that the oxylipin pathway influences aroma, tastes, and maybe deterioration of fresh plant products [81,85]. However, its significance in plant physiology was very incompletely understood until about 60 years later. In the last few years, novel possible physiologic roles for the oxylipin pathway have been accumulating quite rapidly, especially after the demonstration of jasmonins as a chemical messenger in plant stress responses. These discoveries have resulted in new applications of LOX pathway products to the preservation of fresh plant produce during postharvest storage. For example, C\textsubscript{6} aldehydes produced by the LOX/HPLS pathway are demonstrated to be effective fungicides, bactericides, and insecticides, and are used by plants in pest defense [86,87]. Several research groups are now testing the application of C\textsubscript{6} aldehydes to control microbial growth on fruits after harvest. Since C\textsubscript{6} aldehydes are natural compounds in plant tissues, this treatment is considered to be much more acceptable than the use of synthetic biocides. Jasmonic acid, a product of the LOX/HPDS (also known as ADS) pathway, is found to function in plants as a chemical message that triggers protective responses to stresses. Jasmonin treatment of chilling-sensitive fruits and vegetables reduces chilling injury and is suggested for storage of these commodities [88–91].

Thus, in addition to introducing the basic concepts of the oxylipin pathway, this section also focuses on some recent advances into our understanding of its physiological significance in plants. Study of this mechanism should result in new insights into food aspects of the oxylipin pathway product functions. Since the oxylipin pathway greatly interests plant physiologists and biochemists, the next several years should be fascinating times for research in this field.

1. Lipoxygenase (LOX)

The term “lipoxygenase” (linoleate: oxygen oxidoreductase, EC 1.13.11.12) refers to a group of enzymes that catalyze the dioxygenation, by molecular oxygen, of PUFAs containing Z,Z-pentadiene moieties; the resulting enantiomeric hydroperoxy fatty acids have the Z,E-diene conjugation. Enzymes of this type were formerly known as fat oxidase, carotene oxidase, and lipoxidase and have also been listed under EC 1.99.2.1 and EC 1.13.1.13.

LOXs are widespread throughout the plant kingdom [92], being found among the algae, bryophytes, and vascular plants. Considerable LOX activity is detected in various plant organs, such as seeds, stems, leaves, fruits, tuber, and florets [92,93].
In general, young and developing plant tissues or organs have higher LOX activity than the mature ones [94,95]. In cells, LOX is primarily a soluble, cytoplasmic enzyme. LOX is also associated with the chloroplasts in photosynthetic tissues [96], vacuoles in paraveinal mesophyll cells [97,98], and lipid bodies in oilseeds [99].

Many plants contain multiple LOX isozymes encoded by different genes. It is estimated that there are up to 12 different LOX genes expressed as different developmental stages in the life cycle of the soybean [100,101]. Peas have at least seven isoforms, five of which are minor [102]. The isoforms show different properties and likely play different roles throughout the life cycle of plants [99]. During soyseed germination, the seed LOX isoforms disappear, with increases in synthesis of the seeding iso-LOXs [103,104].

Soybean seed LOX was one of the first enzymes to be purified and crystallized [105] and is the best characterized of all plant LOXs. It is composed of three major isoforms, L-1, L-2, and L-3, and one minor isoform, L-4. L-1, L-2, and L-3 show differences in regiospecificity, pH optimum, and enzymatic properties. The type 1 LOX (L-1) catalyzes the incorporation of oxygen predominantly at C-13 of 18:2 and 18:3 and has its optimum activity at pH 9. Its preferred substrate is free fatty acid. Type 2, or mixture LOXs (L-2 and L-3), which have their maximum activities at neutral pH values, do not show positional selectivity for oxygen incorporation. Indeed, they catalyze the formation of approximately equal proportions of 9- and 13-hydroperoxide isomers. Preliminary studies with purified isozymes indicate that L-2 mostly produces the 13-hydroperoxide and L-3 the 9-hydroperoxide (unpublished results). L-2 and L-3 are about equally active with 18:2, methyl 18:2, and linoleoyl glycerol [106,127].

All three major soyseed LOXs are composed of single polypeptide chains of similar lengths (about 96 kDa and 850 amino acids). The complete sequence of the LOX isoforms [108–110] shows that L-1, L-2, and L-3 share high homology in the carboxyl-terminal half of their polypeptide chains. Within this section, there is a region known as the “histidine region,” which is particularly highly conserved in both plant and animal LOXs [111–113]. Missense mutation substituting glutamine for histidine—532 result in the loss of LOX2 activity and protein from mature soybean seeds [113a]. Sequences of LOXs from a variety of plants have led to classification of LOXs into two major groups, LOX1 and LOX2, based on general homologies [114]. Most plant LOXs are of the LOX1 class, having DNA sequence like soybean L-1. The LOX2 class has putative plastid transit sequences, indicating that they are chloroplastic enzymes.

LOXs contain one atom of nonheme iron. The iron atom in lipoxygenase alternates between the Fe(II) and the Fe(III) states during catalysis. The native-resting LOX, a relatively inactive form, is in the high-spin Fe(II) state. Activation of the native E-Fe(II) requires oxidation of the iron atom from Fe(II) to Fe(III) by the reaction product, fatty acid hydroperoxides or H₂O₂ [86]. Because of the product activation requirement, the oxygenation reaction exhibits a characteristic initial lag period.

The LOX-catalyzed reaction is initiated by the stereospecific removal of hydrogen from the C-11 methylene group when 18:2 is used as a substrate (Fig. 9). This occurs possibly by a basic amino acid, followed by the transfer of an electron back to the iron atom to reduce it to E-Fe(II).
Figure 9  Proposed scheme for lipoxygenase oxidation of linoleic acid under aerobic and anaerobic conditions. Fatty acid structures are abbreviated to show only C-8 through C-14. (From Ref. 86, courtesy of H. W. Gardner.)

Under aerobic conditions, this hydrogen abstraction results in the free radical of the fatty acid. When the 11-pro(S)-hydrogen (or D-hydrogen with Fischer projection) is removed, the unpaired electron comes to reside on C-13. Oxygen attacks C-13 to form 13(n-6)S (or L)–hydroperoxy-9Z,11E-octadecadienoic acid. However, when the 11-pro(R)-hydrogen (or L-hydrogen with Fischer projection) is removed, the unpaired electron comes to reside on C-9 and oxygen attacks C-9 to form 9(n-10)S (or D)–hydroperoxy-10-E,12-Z-octadecadienoic acid. The Z double bond at the point of attack isomerizes to the E configuration and moves into conjugation with the neighboring Z double bond. The stereospecific removal of a C-11 methylene hydrogen is a rate-limiting step [115].

Molecular oxygen reacts with the linoleoyl radical to form a peroxyl radical, which subsequently accepts an electron from Fe(II) and acquires a proton to complete the hydroperoxidation, forming regiospecific hydroperoxides. Finally, linoleoyl hydroperoxide (HPOD) dissociates from the enzyme, which has returned to its active E-Fe(III) state.

The specific point of oxygen attack (the regiospecificity) of LOXs depends on the sources of the enzyme, isozymes, and substrates, and on the experimental conditions. LOXs of soybean seed and tea leaves catalyze the incorporation of oxygen predominantly at C-13 of 18:2 or 18:3. Tomato (Lycopersicon esculentum Mill.),
potato, and corn have LOXs that oxygenate primarily at C-9. LOX regiospecificity can vary with LOX isozymes. At pH 6.8, pea L-1 produces mainly C-9 HPOD; however, pea L-2 forms predominantly C-13 HPOD [116]. With LOX from flaxseed (*Linum usitatissimum* L.), 80% of the 13-isomer was formed when 18:2 served as substrate and 88% of this isomer was formed when the linolenic acid was substituted for 18:2 [117]. The pH at which the reaction is conducted, the O₂ concentration, and the temperature also influence the 9:13 ratio of the products [118]. Soybean LOX-1 catalyzes the oxidation of 18:2 into both 13S-HPOD and 9S-HPOD at pH values less than 9. However, the negligible percentage of 9S-HPOD is produced at pH higher than 9. Replacement of histidine-608 by valine in amino acid sequence of cucumber lipid body LOX altered the positional specificity of this 13-LOX in favor of 9-lipoxygenation [117a].

The regiospecificity of oxygen addition to the linoleoyl radical also depends on the ability of the enzyme to recognize the carboxyl or methyl terminal end of the substrate. At pH 9, soybean L-1 recognizes and orients the methyl end of the 18:2. The enzyme abstracts the pro(S)-hydrogen from C-11, and catalyzes the formation of 13S-HPOD [119]. LOXs from wheat, potato tubers, and maize kernels [86] recognize the carboxyl group, abstract the R-hydrogen from C-11, and catalyze the formation of 9S-HPOD.

When the oxygen supply is depleted, the activated enzyme abstracts a hydrogen atom from C-11 as usual to form a linoleoyl radical, and the enzyme returns to the native E-Fe(II) form. Because no oxygen atoms are available to react with the radical, it dissociates from the enzyme to form a free radical, which results in the formation of a variety of products such as fatty acid dimers, ketone, and epoxides. The native E-Fe(II) could be reactivated by oxidation with hydroperoxy products. Reduction products of the hydroperoxide during the activation are hydroxyl ions and alkoxy radicals, which rearrange or combine to form oxodienoic acids, dimers, and pentane.

If PUFAs containing more than three double bonds are used as substrates, LOX can catalyze the incorporation of a second oxygen molecule into a fatty acid hydroperoxide. Soybean and potato tuber LOXs convert α-18:3 to 9,16-dihydroperoxy-10E,12Z,14E-octadecatrienoic acid [120,121]. Soy LOX also catalyzes the formation of 8S,15S-dihydroperoxy 5Z,9E,11Z,13E-eicosatetraenoic acid [122] or 5S,15S-dihydroperoxy-6E,8Z,11Z,13E-eicosatetraenoic acid [123], using 20:4 as a substrate.

It is widely accepted that LOX recognizes the (1Z,4Z)-pentadiene moiety. However, unsaturated fatty acids which have C₈–C₁₂ chains do not act as substrates for tea chloroplast LOX even though they possess the (1Z,4Z)-pentadiene moiety in their structure. All the C₁₀-fatty acid acting as substrates had the (1Z,4Z)-pentadiene moiety between C-9 and C-12 positions numbered from the carboxyl group (ninth and sixth from methyl group; α₁₀ and α₉). The geometric isomers of 18:2, (9E,12E)-, (9E,12Z)-, and (9Z,12E)-octadecadienoic acid did not act as substrates. Ten positional isomers of 18:2 of (3Z,6Z)- to (13Z,16Z)-octadecadienoic acid that do not occur naturally [except for the (9Z,12Z)-acid 18:2] are oxygenated by tea chloroplast LOX. However, the rate of oxygen uptake is 30–60% lower than 18:2 [124]. When the entire series of (α₆Z, α₉Z)-C₁₄–C₁₄ dioenoic and (α₃Z, α₆Z, α₉Z)-C₁₄–C₁₄ trienoic acids are used as substrates, soybean LOX1 activity increases from C₁₄ to C₂₀ and decreases thereafter. No appreciable activity was detected with C₁₁ and C₁₅ substrate. When the entire series of C₁₄–C₁₄ dioenoic acid with a fixed (9Z,12Z)-C₁₃–diene carboxyl moiety are used as substrates, LOX activity is minimal compared with the C₁₆, lin-
oleic acid. These results suggest that the substrate requirement for the hydrophilic area of LOX is fairly broad, but in contrast, that for the hydrophobic area is strict [124,125]. In addition to fatty acids, certain glycerides, fatty esters, and other fatty acid derivatives also act as substrates for LOX [81,119,126,127]. However, the substrate specificity obviously depends on LOX isozymes. Soybean LOX1 is relatively more effective in the lipid-dependent O₂ uptake of free fatty acids compared to esterified derivatives, whereas LOX2 or 3 is relatively more effective with esterified 18:2 than LOX1 [127].

One of the oldest but least understood characteristics of LOXs is their ability to catalyze the bleaching of various pigments, such as carotene, xanthophylls, chlorophyll, cholesterol, crocin, luein, and various dyes. This is a cooxidation reaction that has been used as the basis for some LOX assays and in some industrial practices.

Pigment destruction during lipid peroxidation is usually hypothesized to stem from reactive intermediates generated in the peroxidative reaction [128], and the intermediates are thought to be free radicals. The observations that LOX is unable to bleach when only preformed hydroperoxide is added to the reaction mixture and that bleaching is able to occur anaerobically [126,129] indicate that pigment bleaching is a radical-mediated pathway. The reports that the reaction conditions that can retain conjugated diene production selectively inhibit bleaching and that the bleaching is closely coupled with decreased carbonyl compound production (indicative of peroxide breakdown) suggest that compounds produced before fatty hydroperoxide formation are the ones active in bleaching.

Most models for cooxidation reaction involve the dissociation of the enzyme radical complex. Both Weber et al. [130] and Grosch et al. [131] favor the lipid peroxide (LOO⁻) as the reactive species in bleaching. However, Veldink et al. [119] suggest that dissociation produces the unoxygenated fatty acid radical, which is then free to attack other molecules, including pigments. Lack of stereospecificity of the hydroperoxides formed and no requirement of aerobic conditions in bleaching implicate the unoxygenated fatty acid pathway.

The bleaching activity of LOX depends on the enzyme sources and isozymes. Enzymes from peas and beans (Phaseolus sp.) and the L-2 and L-3 from soybean have a high cooxidation potential; potato LOX is intermediate, whereas the cooxidation activity of wheat, flax, and soybean L-1 is poor [131,132]. The observed differences in cooxidizing potentials among LOXs might result from the different strengths of the associative bonds between enzyme and radical.

LOXs are inactivated by LOX substrate analogs and lipid antioxidants. The antioxidants most commonly used as LOX inhibitors are α-tocopherol, nordihydroguaiaretic acid (NDGA), propyl gallate, hydroquinone, and α-naphthol. These compounds are generally thought to inhibit LOX by their ability to scavenge free radicals, to reduce Fe(III), or both. Thus they are not truly specific LOX inhibitors. LOX substrate mimics include competitive inhibitors (e.g., 18:1, PUFAs with E double bonds) and suicide inhibitors, such as 5,8,11,4-eicosatetraynoic acid (ETYA), 12-iodo-9Z-octadecenoic acid, colneleic acid, and hexanal phenylhydrazone. So far, only suicide inhibitors are considered to be specific LOX inhibitors and can be acceptably used as one of the criteria for the identity of true LOXs. In addition, hydroxamic acids, such as salicylhydroxamic acid (SHAM), phenylhydrazine/phenylhydrazones, or nonsteroidal anti-inflammatory drugs, also cause inhibition of LOXs. The mech-
anistic studies of LOX inhibitors were reviewed by Veldink and Vliegenthart [133] in 1991.

LOX is also a self-destructive enzyme; that is, LOX catalyzes its own destruction during oxygenation of PUFA substrates [134,135]. Soybean L-1 undergoes inactivation during incubation with PUFA or the exposure to hydroperoxy acids [135,136]. The inactivation potency is in the following order: 15-hydroperoxy 20:4 > 15 hydroperoxy 20:3 > 9-hydroperoxyoctadecatrienoic acid (HPOT) or 13-HPOT. The velocity of the LOX-catalyzed reaction decreases as a linear function of substrate utilization. This inactivation seems to require a homolytic cleavage of peroxide group and to result from binding of unstable intermediates produced from monohydroperoxy acids that are bound to the active site. The activity of soybean L-1 may thus be regulated by this mechanism between the secondary oxygenation and autoinactivation [137]. LOX self-destruction could also be via modification of protein by lipid hydroperoxide-derivative free radicals and aldehydes [135].

LOX products, fatty acid hydroperoxides, and their free radical derivatives are very reactive compounds that result in disruption of membrane integrity [82,138], inactivation of proteins and amino acids by reacting with sulfhydryl groups [139], and degradation of DNA by hydroperoxide attack on guanine nucleotides [140]. These properties have led to speculation that lipoxygenase participates in the senescence process in plants. Mazliak [141] and Watada et al. [85] assume that LOX plays a key role in the deterioration of plant produce after harvest. In their hypothesis, processing of vegetables and fruits during harvesting or after harvest results in the release of PUFA's from glycerides. Free PUFA's are then oxygenated by LOX to form lipid hydroperoxides. Both lipid peroxidation and lipid hydroperoxides promote loss of membrane integrity, protein, and chlorophyll, causing accelerated deterioration of plant tissues. Changed activity of LOX has been reported as one of the early events in plant senescence. During fruit ripening and storage, LOX activity was shown to rise [142–144]. Pauls and Thompson [145] found that with advancing age of the cotyledons of Phaseolus vulgaris, LOX activity increased with enhanced lipid hydroperoxides in microsomal membranes, suggesting that these increases ultimately lead to membrane disruption in senescing tissues. In senescing bean cotyledons and carnation (Dianthus caryophyllus L.) flowers, Thompson et al. [146] found that increased LOX activities were associated with fluorescent products of lipid peroxidation. With pea leaf senescence, LOX activity increased by more than sixfold. Addition of LOX inhibitors retarded the senescence with reduced activation of LOX activity by about 50% [147,148]. Senescing carnation petals showed phospholipid degradation and a rise in membranous LOX activity [149,150]. Phenidone, a known LOX inhibitor, delayed senescence in carnations [151].

The literature provides contradictory data regarding changes in LOX activities during deterioration. Declines in LOX activity occurred during senescence of wheat and rye leaves [152] and soybean cotyledons [153]. Twenty-month-old potato tubers had a lower activity of LOX than 8-month-old tubers [154]. During broccoli (Brassica oleracea L.) floret deterioration, increases in lipid peroxidation are closely associated with reduced LOX activities [155–157]. In addition, accumulated studies show that higher LOX activity is more commonly associated with young and fast growing rather than senescing plant tissues and organs [158]. For example, LOX activity decreased by 70% during soybean leaf maturation on plants [95]. As the soybean cotyledons turned yellow and senesced, LOX activity decreased. Removal
of the seed pod reversed senescence (the cotyledons rejuvenate) with increased LOX activity [153]. These phenomena evidently are not in agreement with a role for LOX in senescence. It is worth mentioning that these contradictive results are all based on in vitro measurements, and the role of LOXs in developmental processes cannot readily be assessed until more information is available on in vivo LOX product formation in plant tissues under different conditions.

Increased LOX activity in oilseeds during the early stages of germination has recently been presumed to be involved in the initiation of the mobilization of storage lipids. In this hypothesis, Feussner et al. [99] propose that during early stages of cucumber seed germination, a LOX form is induced and is activated by binding to the lipid body membrane. The active enzyme then oxygenates the esterified fatty acids located in the lipid storage organelles to form hydroperoxides, both in the storage triacylglycerols and in the phospholipids of the lipid body monolayer membrane. The oxygenated fatty acids are preferentially cleaved and subsequently released into the cytoplasm. In the cytosol, the hydroperoxide derivatives are utilized via β-oxidation to serve as a major carbon source for the seedling before full photosynthesis is developed. However, studies of direct association of LOX with lipid bodies in germinating soybean seeds in vivo have been negative [159]. LOXs have also been proposed to be involved in nitrogen partitioning and storage in plants. During soybean seed development, members of LOX multigene family, so-called vegetative LOX, function in nitrogen and assimilate partitioning, mechanisms evolved by plants to temporarily store and subsequently remobilize nutrients to meet specific needs. The levels of gene transcript and protein accumulation of vegetable LOXs in mature soybean generally increase in response to increasing levels of available nitrogen within cells and tissues. LOX genes are regulated in response to plant nitrogen status in both a developmental- and tissue-specific manner. Removal of developing pods, a strong assimilate sink, causes a reallocation of nitrogen and other assimilates to vegetative LOX [160–162]. Kolomiets et al. [163] recently reported that the potato LOX1 class of genes is involved in potato tuber enlargement.

It has been suggested that LOX may also have been involved in the formation of biological functional compounds such as phytoalexin, abscisic acid, and ethylene, as well as in the regulation of the Calvin cycle, the response to wounding and pathogen attack [81,94]. Antisense-mediated depletion of a potato LOX reduces wound infection of proteinase inhibitors with increased weight gain of insect pests and tuber yields. However, the regulatory role of LOX is not caused by its involvement in the wound-induced increase of JA, as wild-type and LOX3-deficient plants have similar jasmonate levels after wounding [164].

A more universal role for plant LOX, however, apparently is to provide fatty acid hydroperoxide substrates for several enzyme systems designated as the HPLS pathway, the HPDS (allene oxide synthetase) pathway, the HPIS pathway, and the HPPR/HPEP pathway.

2. Hydroperoxide Metabolism

a. Hydroperoxide Dehydratase Pathway–Jasmonic Acid Biosynthesis. HPDS (EC 4.2.1.92), also called allene oxide synthetase, is the first enzyme of this pathway (Fig. 10) that leads to the biosynthesis of (+)-7-isojasmonic acid, a plant growth regulator. The enzyme catalyzes the dehydration of a fatty acid hydroperoxide to form an allene oxide [165,166]. The allene oxide 12,13S-epoxy-9Z,11E,15Z-octa-
decatrienoic acid is extremely unstable and has a half-life of only 26 seconds at 0°C [167]. Two types of product result from nonenzymatic transformation of the transitory allene oxide: ketols and a cyclopentenyl metabolite. Spontaneous hydrolysis of the allene oxide results in formation of an α-ketol, 12-oxo-13-hydroxy-9Z,15Z-octadecadienoic acid, and a γ-ketol, 12-oxo-9-hydroxy-10E,15Z-octadecadienoic acid. The α-ketol is the predominant product. The second type of product, a cyclopentenyl compound, results from the spontaneous cyclization of the allene oxide to form 8-[2-(Z-2'-pentenyl)-3-oxo-cyclopent-4-enyloctanoic acid (Fig. 11), which is given the common name 12-oxo-phytodienoic acid (12-oxo-PDA). However, the formation of 12-oxo-PDA can also be due to enzymatic catalysis [167].

In plants, only the 13S-HPOT is converted into the cyclic fatty acid, whereas allene oxides from the 9S-HPOT, 9S-HPOD, and 13S-HPOD exclusively undergo hydrolysis to the ketols. Even with the 13S-HPOT, spontaneous hydrolysis of allene oxide to the ketols occurs to a larger extent compared to the cyclization. Spontaneous cyclization causes the formation of racemic 12-oxo-PDA (9S,13S, and 9R,13R); but in the presence of allene oxide cyclase (EC 5.3.99.6), mainly one isomer (98% 9S,13S) is found in increased yield at the expense of ketol formation [167]. The
double bond at C-15 of the allene oxide intermediate appears to facilitate the cyclization reaction.

The next steps in the pathway to jasmonic acid involve the reduction of the double bond of cyclopentenone ring of 12-oxo-PDA followed by three successive \( \beta \)-oxidations [168]. The reduction reaction is catalyzed by 12-oxo-phytodienoate reductase (EC 1.3.1.42), utilizing NADPH as the reductant to produce \((1S,2S,3\text{-oxo-2(2'}\text{-pentenyl})\text{cyclopentaneoctanoic acid (abbreviated OPC-8:0, where 8:0 refers to the eight-carbon side chain). OPC-8:0 retains the same } E\text{ stereochemistry of the side chains as its precursor.}

The existence of \( \beta \)-oxidation enzymes in the \((+7\)-isojasmonic acid pathway has only been inferred on the basis of the observed products, which are OPC-6:0, OPC-4:0, and OPC-2:0 \([(+7\)-isojasmonic acid]; no metabolites with odd-numbered side chains have been identified [169].

Zimmerman [170] was the first to observe that HPDS in flaxseed with tranform 13S-HPOD into 12-oxo-13-hydroxy-9\( ^Z\)-octadecenoic acid (\( \alpha \text{-ketol). This is the work that led to the decription of the enzyme as a hydroperoxide isomerase. Zimmerman
and Feng [171] later found a “new” enzyme that catalyzes the conversion of 13S-HPOT into a cyclic fatty acid, 12-oxo-PDA, and consequently the term “hydroperoxide cyclase” was used for this reaction. In 1987 Hamberg [165] showed that the α- and γ-ketols are derived from hydrolysis of the actual enzymatic product, an allene oxide fatty acid. He renamed the enzyme as hydroperoxide dehydrase. Researchers soon recognized that the allene oxide fatty acid was the intermediate involved in cyclization of 13S-HPOT into 12-oxo-PDA [172]. Based on these findings, in 1991 the nomenclature was changed again to allene oxide synthase [173]. However, the International Union of Biochemistry and Molecular Biology gives this enzyme the name hydroperoxide dehydratase (HPDS) [174].

HPDS occurs in most cereal seeds or seedlings, flaxseed, lettuce (Lactuca sativa L.), sunflower, spinach, cotton seedlings, Vicia faba, eggplant (Solanum melongena L.), and Chlorella pyrenoioclosa [175]. HPDS is a membrane-bound enzyme and routinely isolated as a microsomal pellet. In spinach leaves, both HPDS and HPLS are associated with whole and broken chloroplasts [176].

Study of the molecular biology of jasmonic acid biosynthesis implies that linolenic acid is converted to jasmonic acid by a process that is likely to begin in chloroplasts and end in peroxisomes, at least in leaves. Conversion of linolenic acid to 12-oxophytodienoic acid in chloroplasts is accomplished through a multistep enzymatic process involving lipoxygenase, allene oxide synthase, and allene oxide cyclase. Jasmonic acid synthesis proceeds with the action of a cytoplasmic 12-oxophytodienoic acid reductase and three rounds of β-oxidation that take place in peroxisomes. Most of the enzymes and the corresponding coding genes involved have been isolated and characterized in different plant species. The majority of these genes are transcriptionally activated by wounding and some of them are also activated by JA, allowing feedback regulation of the biosynthetic pathway [177]. Many wounding-inducible genes are induced by JA but only some insect-inducible genes are induced by JA [178]. A mutant of Arabidopsis, which lacks a functional 3-ketolacyl-CoA thiolase protein and is defective in glyoxysomal fatty acid β-oxidation, has been extensively characterized [179]. A comparison of thiolase genes between the wild type and mutant revealed that the ATT codon for Ile100 at the fourth exon in the wild type is changed to ATGG in the mutant. This nucleotide substitution of T to GG causes a frameshift and produces a stop codon within the fourth exon. Therefore, the mutant thiolase gene encodes a smaller protein, which is unstable and is degraded rapidly [179].

Experiments by Meshack Afithile (personal communication) indicate that this thiolase mutant has reduced biosynthesis of JA but not nearly to the extent of the decrease in normal fatty acid β-oxidation, suggesting that there are both similarities and differences between standard peroxisomal fatty acid β-oxidation and JA biosynthesis.

The specificity of HPDS toward the 13- or 9-HPOD or HPOT is species-dependent. In flaxseed and cotton, there is a strong preference for the 13-hydroperoxide isomer [180,181], whereas the enzyme from corn germ shows no preference. The HPDS gene from flaxseed was cloned and sequenced in 1993 [182]. The purification and characterization of the enzyme showed that it is a cytochrome P450 with an M, of 55 kDa [173]. HPDS from other plant sources frequently, however, has an M, between 220 and 250 kDa [96,181], suggesting that it may exist as a tetramer.

An intriguing role proposed for jasmonic acid that has received wide attention is its function as a signal (i.e., a chemical messenger) in response to certain stressors.
Since 1971, when this physiologic role was first suggested by Anderson [183], further evidence has strengthened the hypothesis. For example, water stress (desiccation) induces the accumulation of some characteristic polypeptides in barley leaves. The application of methyl jasmonate to healthy, unstressed leaves results in induction of the same polypeptides [184]. When soybean plants were stressed by three treatments: removal of seed pods, block of phloem export and water deficit, any of these three treatments induces the synthesis of leaf vegetative storage proteins (VSPs), a group of proteins that are thought to be repositories for nitrogen and can be mobilized under conditions favorable for growth and transported to developing plant organs. Application of methyl jasmonate to soybean plants mimics the effects of these treatments by inducing VSP synthesis in healthy soybean plants [85,186]. In transgenic Arabidopsis plants, where cosuppression reduces chloroplast LOX2 accumulation, the wound-induced accumulation of jasmonate is absent, and there is less VSP mRNA formation compared to the control in response to wounding [187]. Gaseous methyl jasmonate promotes tendril coiling in Bryonia dioica Jacq., and the kinetics of the coiling parallels that of coiling induced by mechanical stimulation [188]. Cultured cells of Rauwolfia canescens and Eschscholtzia californica (California poppy) can be induced by a yeast elicitor to synthesize low molecular weight defensive compounds derived from the phenylpropanoid pathway [189]. The elicitor also enhances sharply the concentration of jasmonic acid within 30 minutes after exposure. When jasmonins are applied to cell cultures in the absence of the yeast elicitor, they independently induce the synthesis of defensive compounds. Proteinase inhibitors are low molecular weight proteins and interfere with insect digestion, prompting a decline in feeding by the insect. Synthesis of the inhibitors is activated either by insects chewing on the leaf or by mechanical damage to the leaf. Accumulation of the inhibitors is both local and systemic. Application to the leaf of jasmonic acid, or its biosynthetic precursors, leads to the same specific induction of both proteinase inhibitor mRNA and protein synthesis. In addition, exposure of tomato plants to gaseous methyl jasmonate, or to plant species that synthesize methyl jasmonate (e.g., sagebrush), also stimulates proteinase inhibitor synthesis [190,191]. Treatment of detached tomato leaves with either LOX inhibitors, ETYA and SHAM, or HPDS inhibitor aspirin blocks the wound-induced accumulation of both protein inhibitors and jasmonic acid. The inhibitory effect of aspirin on protein inhibitors can be overcome by addition of exogenous 12-oxo PDA or jasmonic acid [192].

Based on the above observations, a jasmonate cascade was suggested in higher plants [115]. However, understanding the physiological roles of jasmonic acid in these many different processes is complicated by the observation that intermediates in the synthesis of jasmonic acid are also active in at least some of these processes. For example, 12-oxo-PDA has been shown to be a far more effective inducer of tendril coiling in Bryonia dioica than jasmonic acid itself [193,194] and to act as an elicitor of alkaloid biosynthesis in Eschscholtzia californica cell cultures [195]. The levels of jasmonic acid, 12-oxo-PDA, and other intermediates of oxylipin synthesis vary considerably among species [196], giving rise to the suggestion that the relative and absolute concentrations of different compounds of the “oxylipin signature” may provide flexibility to this multifunctional octodecanoid signaling system [197]. Koch et al. [198] found that early, intermediate, and late compounds of the oxylipin cascade leading to JA have differential effects on the induction of volatile biosynthesis.
Other biological effects of methyl jasmonate/jasmonic acid are summarized in Table 1 [88–91,199–202].

b. Hydroperoxide Lyase Pathway. HPLS is an enzyme that cleaves fatty acid hydroperoxides into two fragments at either side of hydroperoxy group to form an oxo-acid and alkane/alkene or their derivatives.

In the higher plants, HPLS cleaves fatty acid hydroperoxides at the position between the hydroperoxide-bearing carbon and the double bond (the so-called heterolytic HPLS pathway). This cleavage results in aldehydic functions on both sides of the site of cleavage, producing both an alkanal/alkenal and an oxo-acid (Fig. 11).

When 13-HPOD and HPOT are used as substrates, HPLS produces hexanal and 3Z-hexenal, respectively, as well as 12-oxo-Z-9-dodecenoic acid. The 3Z-hexenal and 12-oxo-9Z-dodecenoic acid are often isomerized enzymically or nonenzymically into 2E-hexenal and 12-oxo-10E-dodecenoic acid, respectively. The aldehydes could be reduced by alcohol dehydrogenases into the corresponding C₆-alcohols [93,203,204]. Recently, the pathway has been expanded to include the conversion of 3Z-hexenal into 4-hydroxy-2E-hexenal [205,206]. Pulse-chase labeling of carnation petals revealed that C₆-aldehydes and their derivatives were formed within a hydrophobic subcompartment of lipid-protein particles within the cytosol and then presumably are released to the cytosol by blebbing of lipid-protein particles [206]. It is now well established that the C₆-aldehydes produced by HPLS are important components of the characteristic aromas and flavors of many fruits and vegetables. Hexanal has a rancid green odor, whereas 3Z-hexenal and 2E-hexenal have intense grassy and spicy-green odors, respectively. The corresponding alcohols lead to similar, more subdued odors. The C₆-aldehydes and their derivatives have been demonstrated to be responsible for “fresh note” of tomato, apple [207] and banana (Musa spp.); however, higher levels of hexanal result in the rancid off-flavors associated with raw legumes and deteriorated vegetable oils and nut products [81]. We found that the control of C₆-aldehyde synthesis is very complicated in plant tissues. In soybean seeds, LOX isomer 2 stimulated, but L-3 inhibited, C₆-aldehyde production compared to L-1 [127,208]. In Arabidopsis leaves, changes of C₆-aldehyde generation paralleled with alteration of only C₁₈ PUFA composition of chloroplastic lipids [209]. However, no such relationship was found in the soybean leaves and fruit during development [95,210]. An alcohol dehydrogenase (ADH) mutant of Arabidopsis ecotype, which was deficient in ADH activity, resulted in significantly quantitative and qualitative changes in the accumulation of C₆ volatiles compared to the wild-type plants. The total quantity of LOX-derived volatiles was greater on a fresh weight basis in the ADH mutant. Qualitatively, hexanol and 3-hexenol levels were approximately 62% and 51% lower in the mutant, respectively, whereas levels of hexenal were approximately equal to 10 times higher. Hexanal accumulation, however, was unaffected [211].

Cleavage of 9-hydroperoxides proceeds by the process described above, except the products are either 3Z-nonenal or 3Z,6Z-nonadienal from the HPOD or HPOT, respectively, as well as 9-oxo-nonanoic acid (Fig. 11). The 9-hydroperoxide-specific HPLS has been separated from the 13-hydroperoxide-specific lyase, indicating that they are individual isozymes in cucumber [212]. After the C₆-aldehydes have formed, other transformations occur similar to those discussed above for the aldehydes derived from 13-hydroperoxides. The cucumber odor is principally due to 3Z-nonenal and 3Z,6Z-nonadienal.
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<td>Broccoli (flower buds)</td>
<td>Unpublished data</td>
</tr>
<tr>
<td>Chilling tolerance</td>
<td>Zucchini, rice, grapefruit, bell pepper,</td>
<td>88–91</td>
</tr>
<tr>
<td></td>
<td>avocado</td>
<td></td>
</tr>
<tr>
<td>Ethylene stimulation</td>
<td>Tomato, apple (fruits)</td>
<td>199</td>
</tr>
<tr>
<td>Germination inhibition</td>
<td>Oat, wheat, other (seeds), camellia (pollen)</td>
<td>199</td>
</tr>
<tr>
<td>Leaf opening inhibition</td>
<td><em>Mimosa pudica</em> L. (leaves)</td>
<td>199</td>
</tr>
<tr>
<td>Lycopene inhibition</td>
<td>Tomato (fruits)</td>
<td>199</td>
</tr>
<tr>
<td>Tendril coiling</td>
<td><em>Bryonia dioica</em> Jacq. (tendrils)</td>
<td>199</td>
</tr>
<tr>
<td>Volatile ester inhibition</td>
<td>Apple (fruit)</td>
<td>199</td>
</tr>
</tbody>
</table>


Source: Ref. 199, courtesy of H. W. Gardner.
Another type of HPLS (the so-called homolytic HPLS pathway), affording cleavage fragments different from those discussed above, has been reported in fungi, algae, and a species of grass [86]. In algae, the enzyme cleaves the 13-hydroperoxide of 18:2 or 18:3 to produce 5- and 13-carbon fragments, rather than 6- and 12-carbon fragments as observed in higher plants. The freshwater alga *Chlorella pyrenoidosa* possesses an HPLS that converts the 13-HPOD or HPOT to pentane or pentene and 13-oxo-9Z,11E-tridecadienoic acid [213]. HPLS from blue-green alga *Oscillatoria* sp. produces pentanol rather than pentane [214]. In mushroom (*Psalliota bispora* Lange), a LOX may exist that is specific for oxidizing the 10S position of fatty acid. Mushroom HPLS cleaved the 10S-HPOD into 1-octen-3-ol and 10-oxo-8E-decenoic acid [215]. 1-Octen-3-ol is an important flavor component of mushrooms.

The existence of HPLS was first predicted from studies on the biogenesis of volatiles by bananas and other fruits [216]. Since then, the enzyme activity has widely detected among the higher plants and variety of plant organs [93]. Most HPLSs appear to be membrane-bound. In the flesh of cucumber fruit, HPLS is associated with three membrane fractions: plasma membrane, Golgi apparatus, and endoplasmic reticulum [83,217]. The enzyme in leaves is typically found in chloroplasts, where it is bound to the thylakoid membrane [96,218].

HPLSs have been partially purified and characterized from several plant sources, mostly dicotyledons [124,218–221]. Native soybean HPLS, estimated by gel filtration, is approximately 250 kDa [202], but sodium dodecyl sulfate gel electrophoresis of the purified enzyme reveals a protein band with an *M* value of only 62 kDa, suggesting that soybean HPLS exists as a tetramer in its native state. However, the enzyme from tea leaves is probably monomeric in its native state (*M* 63 kDa) [93]. In immature fruits of green bell pepper [222], HPLS is a homotrimer of 55-kDa subunits. The activity is considerably inhibited by lipophilic antioxidants. The enzyme shows strict substrate specificity, and 13-HPOD seems to be a sole substrate in vivo. Tissue-print immunoblot analysis indicated that HPLS was most abundant in the outer parenchymal cells of the pericarp of the fruits. Spectrophotometric analysis of HPLS indicated that it is a heme protein. Although CO treatment of the enzyme causes no appearance of a peak at 450 nm, the spectroscopic feature highly resembled that of HPDS. Further analysis of amino acid sequence showed highest homology with flax HPDS, suggesting that the HPLS gene is a member of a novel family of cytochrome P450 that acts on fatty acid hydroperoxides.

Both 13- and 9-hydroperoxides of C18-PUFAs serve as substrates for the enzyme, but HPLS specificities depend on the enzyme isoforms and 18:2 or 18:3 [83]. The enzyme in nonphotosynthetic plant tissues is typically more active with HPOD, whereas in photosynthetic organs HPLS usually shows higher activity with the hydroperoxides of linolenic acid [93,95,127,223]. In the majority of plants, HPLS is specific for 13-hydroperoxides. However, the enzyme from cucumber and kidney bean leaves utilizes both 9- and 13-hydroperoxide isomers. In pear fruit, only the 9-hydroperoxide isomer is a suitable substrate for HPLS [83].

Studies of substrate specificity of tea leaf HPLS using an entire series of ω-6S hydroperoxy-C14–C24 dienoic and trienoic acids as substrates show that the 22-carbon hydroperoxides but not the natural substrates (i.e., those of 18 C atoms) have the highest reactivities. The reactivities of the hydroperoxides of trienoic acids are always four to ten times higher than those of the dienoic acids [125]. In the soybean seeds, C8-aldehydes are formed in the presence of free PUFA's rather than the PUFA glyco-
erides, although PUFAs in both forms appear to be oxygenated by soy L-2 [107]. In vivo, alteration of C_{16} PUFA composition does not impact C_{6}-aldehyde production by *Arabidopsis* leaves [209]. Although various HPLS enzymes have not been investigated completely in regard to their stereoselectivity for hydroperoxide substrates, a few have been identified as specific for hydroperoxides with the S configuration. Thus far, none has been identified as R-specific.

HPLS-catalyzed reactions in plant tissues are commonly triggered by wounding, resulting in characteristic plant odors. The C_{6}-aldehydes formed by HPLS from C_{16}-PUFAs have been demonstrated in vitro to be effective biocides, with α,β-ununsaturated hexenal showing considerably more inhibitory action [86]. The other cleavage product, 12-oxo-10E-dodecenoic acid, is the so-called wound hormone or traumatin [224]. It stimulates cell division near the wound site, resulting in the formation of a protective callus around the wound. Thus, HPLS may provide two active defense agents in the wounding response of plant tissues: (1) a short chain aldehyde to destroy or inhibit the attacking pests and (2) a wound-healing agent to protect and mend the damaged tissue. The 9-carbon aldehydes resulting from cleavage of 9-hydroperoxy fatty acids probably have similar functions; however, no role has yet been proposed for 9-oxo-nonanoic acid.

The first in vivo evidence of C_{6}-aldehyde-involved plant defense was supplied by Croft et al. [87]. These investigators noted that the inoculation of *Phaseolus vulgaris* L. leaves with an avirulent strain of *Pseudomonas syringae* pv. *phaseolicola* Burkholdera resulted in a burst of hexenals and hexenols. However, a virulent, compatible strain of *P. syringae* pv. *phaseolicola* did not elicit the burst. The burst occurred after 15 to 24 hours, which corresponds to the time of the hypersensitive response of plant tissues to the pest invasion. An in vitro experiment showed that one of the hexenals, E2-hexenal, was particularly inhibitory to the growth of *P. syringae* pv. *phaseolicola*. Because the hexenal/hexenol burst occurred before phytoalexin accumulation, these compounds were hypothesized to be early volatile phytoalexins important to early plant responses.

Aerial treatment of *Arabidopsis* seedlings with 10 μM concentrations of trans-2-hexenal for 4 or 24 hours induced several genes known to be involved in plants defense response, including phenylpropanoid-related genes as well as genes of the LOX pathway. trans-2-Hexenal induction thus closely mimicked the group of genes induced by methyl jasmonate. However, trans-2-hexenal exerted a moderate inhibitory effect on root length relative to similar concentrations of methyl jasmonate and was approximately 10-fold less effective than methyl jasmonate at inducing anthocyanin accumulation in *Arabidopsis* seedlings. It is suggested that C_{6} volatiles of the LOX pathway may act as a wound signal in plants but result in a moderate plant response relative to MJ at both the physiological and molecular level [225].

In addition, Vaughn and Gardner [226] found that 4-hydroxy-E2-nonenal has high fungitoxicity. In animal studies, the 4-hydroxyalkenals had other physiological effects [227] including modulation of gene expression and activation of phospholipase and glucose-6-phosphate dehydrogenase.

HPLS has also been proposed to participate in the oxidative catabolism of triacylglycerols during the carbon mobilization in germinating seedlings [228] and regulation of cellular levels of fatty acid hydroperoxide in plant tissues. Griffiths et al. [229] found that fatty acid hydroperoxide levels were within the range of 0.6–1.7% relative to the total fatty acid content of plant tissues for both photosynthetic
and nonphotosynthetic tissues. While the fatty acid hydroperoxide levels were similar to wild type in the individual LOX antisensed plants with large reductions in LOX, basal fatty acid hydroperoxide levels, by contrast, were elevated by 38% in transgenic potato leaves antisense repression of HPLS.

c. Hydroperoxide Isomerase Pathway. HPIS catalyzes the intramolecular oxygenation of hydroperoxides of fatty acids to form epoxyhydroxy or dihydroxy fatty acids. This enzyme appears to be totally different from the “hydroperoxide isomerase” that was recently renamed HPDS. HPISs were reported in cereal flours during 1970s [86]. Recently Hamberg [230–232] described an enzyme system, in the fungus *Saprolegnia parasitica*, that converts PUFAs into allylic epoxy alcohols. Two enzyme activities were identified to be responsible for the transformation. The first enzyme, α6-LOX, catalyzes oxygenation of 18:2 to 13S-HPOD. The other enzyme, HPIS, converts the 13S-HPOD into a mixture of the two epoxy alcohols: 11R,12R-epoxy-13S-hydroxy-9Z-octadecadienoic acid and 9R,10R-epoxy-13S-hydroxy-11E-octadecadienoic acid. The enzyme system favored the E,Z-conjugated 9S- and 13S-HPOD over the corresponding E,E-conjugated isomers and also used 20:4 and 5,8,11,14,17-eicosapentaenoic acids as substrates. The C20-epoxy alcohols could be subsequently hydrolyzed to produce four isomeric trihydroxy-eicosatrienoic acids [86]. In addition, this HPIS catalyzes slow isomerization of unnatural 13R-HPOD.

HPIS is involved in diol fatty acid biosynthesis by red alga (*Gracilariopsis lemaneiformis*). Jiang and Gerwick [233] found that an acetone powder preparation of red algae catalyzed the conversion of 20:4 to 12R,13S-dihydroxy-5Z,8Z,10E,14Z-eicosatetraenoic acid. Further studies reveal that conversion of 20:4 occurs by initial 12-LOX-catalyzed oxygenation into 12S-hydroperoxy eicosatetraenoic acid followed by the HPIS-catalyzed conversion of the hydroperoxide into 12R,13S-dihydroxy-5Z,8Z,10E,14Z-eicosatetraenoic acid [234]. This HPIS converts C18 fatty acid hydroperoxides (e.g., 13S-HPOD, 13R-HPOD, 9S-HPOD, 13S-HPOT) into corresponding diol fatty acids as well.

The fungus *Gaeumannomyces graminis* also contains HPIS, which catalyzes formation of another diol fatty acid, 7S,8S-dihydroxy-9Z,12Z-octadecadienoic acid, following incubation with 18:2 [235]. Linoleic acid is initially oxygenated into 8R-hydroperoxy-9Z,12Z-octadecadienoic acid in the presence of the dioxygenase called 8R-dioxygenase. The resulting hydroperoxide is further converted to 7S,8S-dihydroxy-9Z,12Z-octadecadienoic acid [79] by action of a HPIS. Oxygen isotope (18O) studies showed that the HPIS-catalyzed conversion occurred by an intramolecular process: that is, the terminal hydroperoxide oxygen was utilized by the isomerase for hydroxylation of the C-7 methylene group.

Steric analysis shows that the configurations of the epoxide groups of the α,β- and γ,ε-epoxy alcohols produced from a number of different hydroperoxides are invariably R,R and S,R, respectively, and diol fatty acids produced from several fatty acid hydroperoxide isomers always have the S configuration. Thus, the stereochemistry of the intramolecular oxygenation is dictated by the isomerase [232,234].

d. Divinyl Ether Formation. During the early 1970s, Galliard and coworkers reported that a potato extract catalyzed the transformation of either 9S-HPOD and HPOT to divinyl ether derivatives [9-(1′R,3′R,6Z-nonatrienyl)8E-nonenio acid or 9-(1′E,3′Z,6Z-nonatrienyl)8E-nonenio acid, respectively]. These compounds were named colneleic and colnelenic acids [236]. The enzyme was specific for the
9-hydroperoxides and did not attack the 13-hydroperoxide isomers. The $^{18}$O$_2$ label from $^{18}$O$_2$-labeled 9S-HPOD was inserted into the ether oxygen of the divinyl ether [86]. These unusual ethers represented major lipids in homogenates of potato tubers at pH values exceeding 6.5 but are not present in intact tissue. Colneleic and colnelenic acids could be degraded by oxidative cleavage of the ether to form aldehydic fragments. The degradation was catalyzed by an enzyme in potato tubers but also occurred in the presence of Fe$^{2+}$ ions and some nonheme proteins.

The formation of divinyl ether is catalyzed by soybean LOX in vitro, and knowledge of the mechanism has been used to synthesize divinyl ether from 9S-HPOD by a biomimetic method [86].

3. Hydroperoxide-Dependent Metabolism: Peroxygenase/Epoxygenase Pathways

HPPR or HPEP is a membrane-associated oxidase that catalyzes the epoxygenation of mono- and polyunsaturated fatty acids but only in the presence of fatty acid hydroperoxides. In this pathway (Fig. 12), PUFAs such as 18:2 are first converted to 13S-HPOD by action of LOX. HPPR or HPEP then catalyzes subsequent intermolecular transfer of hydroperoxide oxygen from 13S-HPOD to another unsaturated

![Figure 12](image)

**Figure 12** Biosynthesis of epoxy acids by hydroperoxide-dependent peroxygenase/epoxygenase pathway of linoleic acid in plant tissues: 13S-HPOD, 13S-hydroperoxy octadecadienoate. (Adapted from Ref. 79.)
fatty acid to provide an epoxy fatty acid. As a result, the hydroperoxide itself is transformed into the corresponding alcohol.

In 1977 Ishimaru and Yamazaki [237] reported that a hemoenzyme from pea microsomes catalyzed the hydroxylation of phenols by utilizing HPOD as the oxidant. They named the enzyme peroxygenase; the hydroperoxide itself was converted into the corresponding alcohol. Blee and her colleagues [238,239] showed that soybean microsomes also contain a similar enzyme, sulfoxigenase, which catalyzes the oxidation of sulfides to sulfoxides with a corresponding conversion of fatty acid hydroperoxide into its alcohol. By use of $^{18}$O$_2$-labeled 13S-HPOD, it was determined that the sulfoxide oxygen originates from hydroperoxide [238]. Further investigation disclosed that both microsomal HPPR from soybean [240] and microsomal HPEP from *Vicia faba* [241] catalyze the epoxidation of unsaturated fatty acids in the presence of 13S-HPOD. There is a mole-to-mole epoxidation of olefin to hydroperoxide reduction, and the epoxide acquires one $^{18}$O label from $^{18}$O$_2$-labeled 13S-HPOD. In this work, moreover, the soybean HPPR and sulfoxygenase activities coeluted by CM-Sepharose chromatography, indicating that the same enzyme was involved [242].

Both the soybean HPPR and the *V. faba* HPEP also catalyze the formation of epoxide alcohols. The reduced hydroperoxide, 13S-hydroxy-9Z,11E-octadecadienoic acid, was epoxidized by HPEP to 9S,10R-epoxy-13S-hydroxy-11E-octadecenoic acid and a minor amount of its enantiomeric 9R,10S-epoxide [241]. Although the predominant enantiomer of this isomeric pair was identical to a product of HPIS, the mechanism of its formation was decidedly different. By the use of mixtures of $^{16}$O$_2$- and $^{18}$O$_2$-labeled 13S-HPOD, it was shown that O-transfer to epoxide occurred by both inter- and intramolecular pathways from hydroperoxide [241], in contrast to the strict intramolecular O-transfer by HPIS.

Although HPPR and HPEP are apparently the same enzyme, a few differences between the broad bean HPEP and the soybean HPPR were noted.

1. Both soybean HPPR and *V. faba* HPEP converted 18:1 into its 9,10-epoxide, and 18:2 afforded 9,10- and 12,13-epoxides [220–222]. However, only soybean HPPR could transform 18:2 into the 9,10,12,13-diepoxide [242].

2. The two enzymes produced 12,13-epoxy-9Z-octadecenoic acid of opposite stereochemistry. With *V. faba* HPEP, the 9R,10S and 12S,13R enantiomers predominated over 9S,10R and 12R,13S [221], but with soybean HPPR there was an enantiomeric excess of 9R,10S and 12R,13S over 9S,10R and 12S,13R [242]. That is, the 9,10-epoxides formed by the two enzymes have the same configuration, but the 12,13-epoxides have the opposite stereochemistry.

3. The rates of epoxidation by broad bean epoxygenase of various Z-mono-unsaturated fatty acids were affected more by the position of the double bond than by the length of the carbon chain [243], while the opposite was true for soybean peroxygenase [240].

The peroxygenase pathway may be involved in plant defense at several levels. The peroxygenase pathway is part of the 18:1 cascade involved in the biosynthesis of cutin monomers [244]. The major cutin component, 9,10-epoxy-18-hydroxy stearic acid, and some minor epoxy and/or hydroxy fatty acids are formed by this pathway in a cell free system. Cutin is the chief component of cuticle, which covers all the...
aerial parts of plants, constitutes the first barrier against invasion by pathogens, and also limits the penetration of compounds such as pesticides and herbicides. When pathogens succeed in crossing the cuticle, the plant copes with such aggression by an array of defense mechanisms and, especially, by triggering the synthesis of toxic compounds. The pathway involving hydroperoxide-dependent epoxidation and the secondary conversion of epoxy acids into vicinal dihydroxy acids in the presence of epoxide hydrolase may have significance in plants for protection against fungi and other pathogens [245]. In addition, epoxy fatty acid derivatives may work as an elicitor to induce resistance to pathogens in rice (Oryza sativa L.) [246].

B. Autoxidation

Autoxidation of PUFAs has been demonstrated to result in aging and disorders of animal tissues [247,248] and in meat deterioration [249]. However, it had not drawn much attention from plant scientists until recently, since high activity of LOX is widely distributed in plants. Autoxidation of PUFAs has been suggested to be responsible for deterioration of postharvest plant commodities [248; unpublished results] and senescence of plant tissues [250]. Accumulated data show that the antioxidant defense system is involved in stress responses of plant tissues [251].

Trihydroxyoctadecenoic and hydroxyoctadecadienoic acids in oxidized phosphatidylcholine are responsible for bitterness in soy products and in water suspensions of oat (Avena sativa L.) flour. These bitter-tasting hydroxylated fatty acids are proposed to originate from the free radical decomposition of HPOD [217].

REFERENCES


107. Reference deleted.


I. INTRODUCTION

Autoxidation is a natural process that takes place between molecular oxygen and unsaturated fatty acids. Autoxidation of unsaturated fatty acids occurs via a free radical chain mechanism consisting of basic steps of initiation [Eq. (1)], propagation [Eqs. (2) to (3)], and termination [Eqs. (4) to (6)]. Initiation starts with the abstraction of a hydrogen atom adjacent to a double bond in a fatty acid (RH) molecule/moiety, and this may be catalyzed by light, heat, or metal ions to form a free radical. The resultant alkyl free radical (R') reacts with atmospheric oxygen to form an unstable peroxy free radical, which may in turn abstract a hydrogen atom from another unsaturated fatty acid to form a hydroperoxide (ROOH) and a new alkyl free radical. The new alkyl free radical initiates further oxidation and contributes to the chain reaction, the chain reaction (or propagation) may be terminated by formation of nonradical products resulting from combination of two radical species.

Initiation:

\[ \text{RH} \xrightarrow{\text{initiator}} \text{R}^\cdot + \text{H}^\cdot \]  

(1)
Propagation:

\[ R' + O_2 \rightarrow ROO' \]  \hspace{1cm} (2)

\[ ROO' + RH \rightarrow ROOH + R' \]  \hspace{1cm} (3)

Termination:

\[ R' + R' \rightarrow RR \]  \hspace{1cm} (4)

\[ R' + ROO' \rightarrow ROOR \]  \hspace{1cm} (5)

\[ ROO' + ROO' \rightarrow ROOR + O_2 \]  \hspace{1cm} (6)

The mechanism of lipid autoxidation has been postulated by Farmer et al. [1], Boland and Gee [2], and Bateman et al. [3]. The propagation step in the autoxidation process includes an induction period when hydroperoxide formation is minimal [4,5]. The rate of oxidation of fatty acids increases in relation to their degree of unsaturation. The relative rate of autoxidation of oleate, linoleate, and linolenate is in the order of 1:40–50:100 on the basis of oxygen uptake and 1:12:25 on the basis of peroxide formation [6]. Therefore, oils that contain relatively high proportions of polyunsaturated fatty acid (PUFA) may experience stability problems. The breakdown products of hydroperoxides, such as alcohols, aldehydes, ketones, and hydrocarbons, generally possess offensive off-flavors. These compounds may also interact with other food components and change their functional and nutritional properties [7].

II. MEASUREMENT OF OXIDATIVE RANCIDITY

There are various methods available for measurement of lipid oxidation in foods. Changes in chemical, physical, or organoleptic properties of fats and oils during oxidation may be monitored to assess the extent of lipid oxidation. However, there is no uniform and standard method for detecting all oxidative changes in all food systems. The available methods to monitor lipid oxidation in foods and biological systems may be divided into two groups. The first group measures primary oxidative changes and the second determines secondary changes that occur in each system.

A. Primary Changes

1. Changes in Reactants

Methods that measure primary changes of lipids may be classified as those that quantify loss of reactants (unsaturated fatty acids). Measurement of changes in fatty acid composition is not widely used in assessing lipid oxidation because it may require total lipid extraction from food and subsequent conversion to derivatives suitable for gas chromatographic analysis. Separation of lipids into neutral, glycolipid, phospholipid, and other classes may also be necessary. However, it has been proven that this method serves as a useful technique to identify class of lipids and fatty acids that are involved in the oxidative changes [8,9] and also to assess lipid oxidation induced by different metal complexes that afford a variety of products [10]. On the other hand, changes of fatty acid composition cannot be used in more saturated oils because this indicator reflects only the changes that occur in unsatu-
rated fatty acids during oxidation (11). Therefore, oxidative changes in marine oils and highly unsaturated vegetable oils may be monitored using this indicator. Similarly, changes in iodine value due to loss of unsaturation during accelerated oxidation studies may be used as an index of lipid oxidation [12].

2. Weight Gain

It is generally accepted that addition of oxygen to lipids and formation of hydroperoxides is reasonably quantitative during initial stages of autoxidation. Therefore, the measurement of induction period from weight gain data is theoretically sound. In this method, oil samples (about 2.0 g) are weighed into Petri dishes; then traces of water are removed by placing the samples overnight in a vacuum oven at 35°C and over a desiccant. Samples are then reweighed and stored in an oven at a set temperature. The weight gain of the samples may be recorded at different time intervals.

Olcott and Einset [13] reported that marine oils exhibit a fairly sharp increase in their weight at the end of the induction period and are rancid by the time they gain 0.3–0.5% in weight (at 30–60°C). Ke and Ackman [14] reported that this method is simple, has a satisfactory reproducibility, and may be used to compare oxidation of lipids from different parts of fish. Recently, Wanasundara and Shahidi [15,16] used this method to compare storage stability of vegetable (Fig. 1) and marine oils as affected by added antioxidants and were able to compare relative activity of antioxidants employed. However, surface exposure of the sample to air is an impor-

![Figure 1](image.png)

**Figure 1** Effect of canola extracts (CEs) and commercial antioxidants on the weight gain of canola oil stored at 65°C. BHA, butylated hydroxyanisole; BHT, butylated hydroxytoluene; MGC, monoacylglycerol citrate; TBHQ, tert-butylhydroquinone.
tant variable in determining the rate of oxidation. Therefore, use of equal size containers to store samples is essential when carrying out such experiments.

The weight gain method also suffers from certain disadvantages: (1) the weighing frequency hinders monitoring of fast kinetics (a higher frequency would involve nocturnal weighing), and low or moderate temperatures require long analysis times for stable samples; (2) discontinuous heating of the sample (which must be cooled before weighing) may give rise to nonreproducible results, so the heating and cooling intervals must be accurately controlled; (3) the method involves intensive human participation; and (4) the working conditions (sample size, shape of container, and temperature) may influence the results. Nevertheless, this method offers advantages, such as low instrumentation cost as well as unlimited capacity and speed for sample processing.

3. Hydroperoxides

In the oxidation of fats and oils, the initial rate of formation of hydroperoxides exceeds their rate of decomposition, but this is reversed at later stages. Therefore, monitoring the amount of hydroperoxides as a function of time indicates whether a lipid is in the growth or decay portion of the hydroperoxide concentration curve. This information can be used as a guide for considering the acceptability of a food product with respect to the extent of product deterioration. By monitoring the induction period before the appearance of hydroperoxides, one can assess the effectiveness of added antioxidants on the stability of a food lipid.

4. Peroxide Value

The classical method for quantitation of hydroperoxides is the determination of peroxide value (PV). The hydroperoxide content, generally referred to as PV, is determined by an iodometric method. This is based on the reduction of the hydroperoxide group (ROOH) with iodide ion (I⁻). The amount of iodine (I₂) liberated is proportional to the concentration of peroxide present. Released I₂ is assessed by titration against a standardized solution of sodium thiosulfate (Na₂S₂O₃) using a starch indicator. Chemical reactions involved in PV determination are given below:

\[
ROOH + 2H^+ + 2KI \rightarrow I_2 + ROH + H_2O + 2K^+
\]

Potential drawbacks of this method are absorption of iodine at unsaturation sites of fatty acids and liberation of iodine from potassium iodide by oxygen present in the solution to be titrated [17]. Results may also be affected by the structure and reactivity of peroxides as well as reaction temperature and time. The iodometric method for determination of PV is applicable to all normal fats and oils, but it is highly empirical and any variation in procedure may affect the results. This method also fails to adequately measure low PV because of difficulties encountered in determination of the titration end point. Therefore, the iodometric titration procedure for measuring PV has been modified in an attempt to increase the sensitivity for determination of low PV. The modification involves the replacement of the titration step with an electrochemical technique in which the liberated iodine is reduced at a platinum electrode maintained at a constant potential. PV ranging from 0.06 to 20 meq/kg have been determined in this manner, but it is essential to deaerate all solutions to prevent further formation of peroxides.
Several other chemical methods have also been suggested for monitoring \( PV \). Colorimetric methods based on the oxidation of \( \text{Fe}^{2+} \) to \( \text{Fe}^{3+} \) and determination of \( \text{Fe}^{3+} \) as ferric thiocyanate, and a 2,6-dichlorophenol-indophenol procedure are reported in the literature [18]. In studies on the oxidation of biological tissues and fluids, measurement of fatty acid hydroperoxides is more common than measurement of their decomposition products. Fatty acid hydroperoxides can be analyzed by high-performance liquid chromatography (HPLC) or their corresponding hydroperoxy acid reduction products may be determined by gas chromatography–mass spectrometry (GC-MS) [19]. Fluorescence methods have also been developed to determine hydroperoxides by allowing them to react with substances such as luminol and dichlorofluorescein, which form fluorescent products [17]. Although determination of peroxide value is common, its usefulness is generally limited to the initial stages of lipid oxidation.

5. Active Oxygen and Oil Stability Index/Rancimat Methods

The Active Oxygen Method (AOM), also referred to as the Swift test of the American Oil Chemists’ Society, is a common accelerated method used for assessing oxidative stability of fats and oils. This method is based on the principle that aging and rancidification of a fat is greatly accelerated by aeration in a tube held at a constant elevated temperature. In this method, air is bubbled through a heated oil at 98–100°C for different time intervals and the PVs are determined. The PVs are then plotted against time and the induction period determined from the graph. Even though this method has been used extensively over the years, its inherent deficiencies and difficulties have also been identified. These include the following: (1) the end point is determined by the amount of peroxides in the oxidized oil; peroxides are unstable and decompose readily to more stable secondary products. (2) During the rapid oxidation phase, the reaction is extremely susceptible to variations in the oxygen supply. Automated versions of the AOM apparatus, known as the Oil Stability Instrument (OSI) and Rancimat, are now available. The Rancimat method uses a commercial apparatus marketed by Metrohm Ltd. (Herisau, Switzerland). The OSI, a computer-assisted instrument developed by Archer Daniels Midland (ADM), is now produced commercially by Omnion Inc. (Rockland, MA). These methods may be considered as automated AOM since both employ the principle of accelerated oxidation. However, the OSI and Rancimat tests measure the changes in conductivity caused by ionic volatile organic acids, mainly formic acid, automatically and continuously, whereas in the AOM, peroxide values are determined. Organic acids are stable oxidation products that are produced when an oil is oxidized by a stream of air bubbled through it. In the OSI and Rancimat methods, oxidation proceeds slowly at first because during the induction period formic acid is released slowly. The end point is selected where the rapid rise in conductance begins. The Rancimat became available in the early 1980s and is capable of running only eight samples simultaneously; however, OSI is capable of running up to 24 samples at the same time. In addition, instruments that monitor the drop in the overhead pressure of an oil during heating might be used. An example of this sort of equipment is the Oxidograph, which is commercially produced by Mikrolab (Aarhus, Denmark). In Oxidograph, a sample of oil or fat is exposed to oxygen or air at elevated temperatures. Heating is done in an aluminum block. As the sample absorbs oxygen the pressure change in the reaction vessel is measured electronically by means of pressure transducers.
The rate of oxygen consumption during the early stages of storage of lipids also provides an ideal parameter for shelf life prediction. Wewala [20] used this method for prediction of shelf life of dried whole milk and found very good correlation between the head space oxygen content and storage time (Fig. 2).

6. Conjugated Dienes

Oxidation of polyunsaturated fatty acids is accompanied by an increase in the ultraviolet absorption of the product. Lipids containing methylene-interrupted dienes or polyenes show a shift in their double-bond position during oxidation due to isomerization and conjugate formation [21]. The resulting conjugated dienes exhibit an intense absorption at 234 nm; similarly conjugated trienes absorb at 268 nm.

Farmer and Sutton [22] indicated that the absorption increase due to the formation of conjugated dienes and trienes is proportional to the uptake of oxygen and formation of peroxides during the early stages of oxidation. St. Angelo et al. [23] studied the autoxidation of peanut butter by measuring the PV and absorption increase at 234 nm due to the formation of conjugated dienes. Shahidi et al. [24] and Wanasundara et al. [25] found that conjugated dienes and PV of marine and vegetable oils correlate well during their oxidation (Fig. 3). These authors concluded that the conjugated diene method may be used as an index of stability of lipids in place of, or in addition to, PV. However, carotenoid-containing oils may give high absorbance values at 234–236 nm due to the presence of double bonds in the conjugated structure of carotenoids. The conjugated diene method is faster than PV determination, is much simpler, does not depend on chemical reactions or color development, and requires a smaller sample size. However, presence of compounds absorbing in the same region may interfere with such determinations.

![Figure 2](image_url)  
Figure 2 Change of the headspace oxygen content of stored dried whole-milk samples.
Parr and Swoboda [26] have described an alternate spectroscopic method to determine lipid oxidation of stored oils. In this assay, hydroperoxides of polyenoic fatty acids as well as hydroxy and carbonyl compounds derived from them are converted to more conjugated chromophores by two chemical reaction steps, namely reduction and then dehydration (Fig. 4). These yield *conjugable oxidation products* (COPs), which are measured and expressed as COP values. The first step of the analytical procedure involves reduction of the carbonyl group by sodium borohydride, which results in the disappearance of the characteristic ultraviolet absorption of carbonyl compounds of oxidized polyenoic fatty acids (oxodienes). The decrease in the absorption at 275 nm is known as *oxodiene value*. The next step of the COP assay involves changes in the spectrum of the reduced compound to its dehydrated counterpart which exhibits absorption maxima at 268 and 301 nm. The sum of these absorbance changes at 268 and 301 nm yields the COP value whereas their relative proportions define the COP ratio. For the calculation of oxodiene and COP results the concentration of the final lipid solution also has to be taken into account.

**B. Secondary Changes**

The primary oxidation products (hydroperoxides) of fats and oils are transitionary intermediates that decompose into various secondary products. Measurement of secondary oxidation products as indices of lipid oxidation is more appropriate since secondary products of oxidation are generally odor-active, whereas primary oxidation products are colorless and flavorless. Secondary oxidation products include aldehydes, ketones, hydrocarbons, and alcohols, among others. The following sections describe common methods used for measuring secondary oxidation products of lipids.
1. 2-Thiobarbituric Acid Value

One of the oldest and most frequently used tests for assessing lipid oxidation in foods and other biological systems is the 2-thiobarbituric acid (TBA) test. The extent of lipid oxidation is reported as the TBA value and is expressed as milligrams of malonaldehyde (MA) equivalents per kilogram sample or as micromoles MA equivalents per gram sample. MA is a relatively minor product of oxidation of polyunsaturated fatty acids that reacts with the TBA reagent to produce a pink complex with an absorption maximum at 530–532 nm [27]. The adduct is formed by condensation of two molecules of TBA with one molecule of MA (Fig. 5). Other products of lipid oxidation, such as 2-alkenals and 2,4-alkadienals, also react with the TBA reagent. However, the exact mechanism of their reaction with the TBA reagent is not well understood. There are several procedures for the determination of TBA values. The TBA test may be performed directly on the sample, its extracts, or

\[ \text{TBA} + \text{MA} \rightarrow \text{TBA-MA adduct} \]

Figure 5  Reaction of 2-thiobarbituric acid (TBA) and malonaldehyde (MA).
distillate. In case of the distillation method, volatile substances are distilled off with steam. Then the distillate is allowed to react with the TBA reagent in an aqueous medium. The advantage of the distillation method is the absence of interfering substances. In the extraction method, TBA-reactive substances (TBARSs) are extracted from food material into an aqueous medium (i.e., aqueous trichloroacetic acid) prior to color development with the TBA reagent. The main disadvantages of both of these methods are long assay time and possibility of artifact formation. In the direct assay method, lipid sample (oil) reacts with the TBA reagent and the absorbance of the colored complex so prepared is recorded. The direct assay method is simple and requires less time for sample preparation.

There are certain limitations when using the TBA test for evaluation of the oxidative state of foods and biological systems because of their chemical complexity. Dugan [28] reported that sucrose and some compounds in wood smoke react with the TBA reagent to give a red color that interferes with the TBA test. Baumgartner et al. [29] also found that a mixture of acetaldehyde and sucrose when subjected to the TBA test produced a 532-nm absorbing pigment identical to that produced by MA and TBA. Modifications of the original TBA test have been reported by Marcuse and Johansson [30], Ke and Woyewoda [31], Robbles-Martinez et al. [32], Pokorny et al. [33], Shahidi et al. [34,35], Thomas and Fumes [36], and Schmedes and Holmer [37]. However, it has been suggested that TBARS values produce an excellent means for evaluating the relative oxidative state of a system as affected by storage condition or process variables [38]. Nonetheless, it is preferable to quantitate the extent of lipid oxidation by a complementary analytical procedure in order to verify the results.

Several attempts have been made to establish a relationship between TBA values and the development of undesirable flavors in fats and oils. It has been shown that flavor threshold values correlate well with the TBA results of vegetable oils, such as those of soybean, cottonseed, corn, safflower [17], and canola [5].

2. Oxirane Value

The oxirane oxygen or epoxide groups are formed during autoxidation of fats and oils. The epoxide content is determined by titrating the oil sample with hydrobromic acid (HBr) in acetic acid and in the presence of crystal violet, to a bluish green end point. This method has been standardized by the American Oil Chemists’ Society in their tentative method (Cd 9-57) [39], but it is not sensitive and lacks specificity. The HBr may also attack \( \alpha,\beta \)-unsaturated carbonyls and conjugated dienals, and the reaction is not quantitative with some trans-epoxides. Fioriti et al. [40] found that picric acid was the best of several acidic chromophores in its reaction with epoxides. Despite a nonquantitative reaction, the product concentration followed Beer’s law. This method has been found to be particularly well suited for the determination of epoxides in heated fats and oils where the oxirane content is often less than 0.1%.

3. \( p \)-Anisidine Value

\( p \)-Anisidine value (\( p \)-AnV) is defined as 100 times the optical density measured at 350 nm in a 1.0-cm cell of a solution containing 1.0 g of oil in 100 mL of a mixture of solvent and reagent, according to the IUPAC method [41]. This method determines the amount of aldehyde (principally 2-alkenals and 2,4-alkadienals) in animal fats and vegetable oils. Aldehydes in an oil react with the \( p \)-anisidine reagent under acidic conditions. The reaction of \( p \)-anisidine with aldehydes affords yellowish products,
as shown in Figure 6. List et al. [42] reported a highly significant correlation between 
\( p\text{-AnV} \) and flavor acceptability scores of salad oils processed from undamaged soybeans.

4. TOTOX Value

The \( p\text{-AnV} \) is often used in the industry in conjunction with PV to calculate the so-called total oxidation or TOTOX value:

\[
\text{TOTOX value} = 2\text{PV} + p\text{-AnV}
\]

The TOTOX value is often considered to have the advantage of combining evidence about the past history of an oil (as reflected in the \( p\text{-AnV} \)) with its present state (as evidenced in the PV). Therefore, determination of TOTOX value has been carried out extensively to estimate oxidative deterioration of food lipids [43]. However, despite its practical advantages, TOTOX value does not have any sound scientific basis because it combines variables with different dimensions. Recently, Wanasundara and Shahidi [44] defined TOTOX_{TBA} as \( 2\text{PV} + \text{TBA} \) since determination of \( p\text{-AnV} \) may not be always feasible.

5. Carbonyls

An alternative approach for monitoring the extent of lipid oxidation in fats and oils is to measure the total or individual volatile carbonyl compounds formed from degradation of hydroperoxides. One of the more reliable methods for total carbonyl analysis is based on the absorbance of the quinoidal ion, a derivative of aldehydes.

Figure 6  Possible reactions between \( p\)-anisidine reagent and malonaldehyde.
and ketones. This ion is formed from the reaction of 2,4-dinitrophenylhydrazine (2,4-DDNH) with an aldehyde or ketone, followed by the reaction of the resulting hydrozones with alkali (Fig. 7), which is then analyzed spectroscopically at a given wavelength. Many variations of this spectroscopic method have been reported [45,46]. Each method offers an alternative solvent, wavelength, or workup to analyze the quinoidal ion.

The analysis of individual carbonyl compounds is another method which has recently gained popularity. Hexanal, one of the major secondary products formed during the oxidation of linoleic or other ω-6 fatty acids in lipid-containing foods [47,48], has been used to follow lipid oxidation. Shahidi and Pegg [47] reported that a linear relationship existed between hexanal content, sensory scores, and TBA numbers of cooked ground pork, whereas St. Angelo et al. [49] established a similar correlation for cooked beef. O'Keefe et al. [50] have used hexanal as an indicator to assess oxidative stability of meat from broiler chickens fed fish meal. Supplementation of high amounts of fish meal to the diet increased the hexanal content of the thigh meat during storage. However, recent studies have shown that during oxidation of marine oils which are rich in polyunsaturated fatty acids of the ω-3 type, large amounts of propanal are formed and that a good correlation exists between the content of propanal and the amount of TBARS in such samples [51,52]. Therefore, it is essential to use appropriate indicators when assessing stability of food lipids. We recommend that hexanal be used when oils under investigation are rich in ω-6 fatty acids while propanal would serve as a reliable indicator when oils high in ω-3 fatty acids are being considered.

6. Hydrocarbons and Fluorescent Products

Studies of oxidized methyl linoleate and soybean oil [53] have revealed that saturated hydrocarbons could be detected when aldehydes are either absent or undetectable. Snyder et al. [54] have reported that ethane, propane, and pentane are predominant short chain hydrocarbons formed through thermal decomposition of soybean oil. Correlations of flavor acceptability scores and pentane content determined by GC

![Figure 7](image)

**Figure 7** Reaction steps in the production of hydrozones from carbonyls and 2,4-dinitrophenylhydrazine.
techniques have been used to assess rancidity of fats and oils [55]. Significant correlations existed between the amount of pentane produced and the number of rancid descriptions of stored vegetable oils [56]. Correlation of headspace pentane concentrations and sensory scores of stored freeze-dried pork samples was reported by Coxon [8].

Another secondary change that occurs during autoxidation of biological systems is the formation of fluorescent products from the reaction of MA with amino compounds such as proteins and nucleotides [57]. This method has been used to determine the extent of lipid oxidation in biological tissues. It has been established that fluorescent compounds with a general structure of 1-amino-3-iminopropane may develop through the reaction of an amino group with carbonyl compounds, mainly MA [58–60] (Fig. 8).

Kikugawa and Beppu [61] reported that the development of fluorescence depends not only on the formation of condensation products between malonaldehyde and free amino groups, but also on the nature of the substituents of the latter compounds. Different excitation and emission maxima were observed for different condensation products. Advantages of the fluorescence method as a means of measuring lipid oxidation have been reported by Dillard and Tappel [58]. This method can detect fluorescent compounds at concentrations as low as parts per billion levels and is found to be 10–100 times more sensitive than the TBA assay.

III. MEASUREMENT OF FRYING FAT DETERIORATION

Deep frying is a popular method for food preparation, especially in fast food restaurants. Although vegetable oils are used primarily as a heat exchange medium for cooking, when used for deep frying, they contribute to the quality of fired products. In the process of deep fat frying, a complex series of chemical reactions take place. These reactions are characterized by a decrease in the total unsaturation content of the fat with a concurrent increase in the amount of free fatty acids, cyclic fatty acids

![Figure 8](image_url)

**Figure 8** Production of fluorescent chromophores from the reaction of lipid oxidation products and amines.
(Fig. 9), foaming, color, viscosity, and formation of polar matter and polymeric compounds. As these reactions proceed, the functional, sensory, and nutritional quality of frying fats change and may reach a point where high-quality foods can no longer be prepared. Therefore, it is essential to determine when the frying fat is no longer usable.

Quality evaluation of frying fats may be carried out in many ways. The first attempt to define a deteriorated frying fat was made by the German Society for Fat Research in 1973. It recommended that “a used frying fat is deteriorated if, without doubt, its odor and taste were unacceptable; or if in case of doubtful sensory assessment, the concentration of petroleum ether–insoluble oxidized fatty acids in it was 0.7% or higher and its smoke point was lower than 170°C; or if the concentration of petroleum ether–insoluble oxidized fatty acids was 1.0% or higher.” Although sensory evaluation of foods is the most important quality assessment, taste evaluations are not practical for routine quality control. It is always preferred to have a quantitative method for which rejection point could be established by sensory means. Peroxide values provide an indication of frying fat quality if they are used in a very specific way. However, peroxides generally decompose at about 150°C and hence at frying temperatures (usually 180–190°C) no accumulation of peroxides occurs. Free fatty acids from frying fats can be determined by direct titration with a standardized base in ethanol. Fritsch [62] has shown that in most deep fat frying operations the amount of free fatty acids produced by hydrolysis is too small to affect the quality.

![Figure 9](image)

**Figure 9** Structures of cyclic fatty acids formed during frying.
of foods. However, industrially, frying oil quality is usually checked by measurement of color and/or free fatty acid content in order to tell an operator when a fat is ending its useful life. The foaming characteristics of used fats would also lead one to the same conclusion.

A quick colorimetric test kit is now available for measuring oil quality [63]. Blumenthal et al. [64] developed a spot test to measure free fatty acids in which drops of used fat are placed on a glass covered with silica gel containing a pH indicator in order to give a three-color test scale of blue, green, and yellow. This may indicate the amount of free fatty acids in a sample. Northern States Instrument Corp. (Lino Lakes, MN) has developed an instrument that measures the dielectric constant of insulating liquids. The instrument is a compact unit, relatively inexpensive, simple to operate, and requires only a few drops of oil for each measurement. For evaluation of frying fats, the instrument must be calibrated first with a fresh oil sample prior to its use in frying operations.

Determination of total polar matter in frying fats appears to be emerging as a reliable method for assessing the useful life of fats and oils subjected to frying and is an official method in Europe. Total polar matter is determined by dissolving the fat in a relatively nonpolar solvent, such as toluene or benzene, and running through a silica gel column that adsorbs the polar compounds. After evaporation of the solvent, the nonpolar fat can be weighed and the total polar matter calculated from the weight difference data or determined directly by their elution from the column with diethyl ether or a mixture of chloroform and methanol. Sebedio et al. [65] illustrated that polar and nonpolar fractions of fried oils can be quantitatively estimated using Iatroscan thin-layer chromatography–flame ionization detection (TLC-FID) system with Chromarod SII. This method requires a very small sample and is much faster than silicic acid column separation.

IV. RECENT DEVELOPMENTS FOR QUANTITATION OF LIPID OXIDATION

A. ESR Spectroscopy

Lipid oxidation in foods and biological systems has conventionally been tested by monitoring either primary or secondary oxidation products. Over the last 20 years or so, advances in pulse radiolysis [66] and electron spin resonance (ESR) [67] techniques have facilitated the detection and study of short-lived free radical intermediates. ESR spectroscopy allows selective detection of free radicals. The technique depends on the absorption of microwave energy (which arises from the promotion of an electron to a higher energy level) when a sample is placed in a variable magnetic field. A major limitation in the detection of free radicals by ESR is the requirement that radical concentrations remain higher than $10^{-8}$ M. Radical lifetimes in solution are very short (<1 msec), and steady-state concentrations generally remain well below $10^{-7}$ M. Several approaches have been developed to overcome this problem, either by enhancing the rate of radical production or by diminishing the rate of its disappearance. These techniques include rapid freezing, lyophilization, or spin trapping [68]. Although application of ESR spectroscopy as a precise method to study lipid oxidation in animal tissues and other biological systems is commonplace, its application to foods is relatively new.
Yen and Duh [69] and Chen and Ho [70] have reported that inhibition of free radical formation by different antioxidants can be measured using very stable free radicals such as 1,1-diphenyl-2-picrylhydrazyl (DPPH). The mechanism of the reaction of antioxidant with DPPH radical is as follows:

\[
(DPPH)^* + HO--R--OH \rightarrow (DPPH):H + HO--R--O^* \\
HO--R--O^* + (DPPH) \rightarrow (DPPH):H + O==R==O
\]

DPPH radical, with a deep violet color, receives a hydrogen atom from the antioxidant and is converted to a colorless molecule. Using this reagent, the free radical scavenging ability of the antioxidant can be determined by spectrophotometric methods.

B. Infrared Spectroscopy

Infrared (IR) spectroscopy has also been used for measurement of rancidity, and it is of particular value in recognition of unusual functional groups and in studies of fatty acids with trans double bonds. Production of hydroperoxides during oxidation of lipids gives rise to an absorption band at about 2.93 μm, whereas the disappearance of a band at 3.20 μm indicates the replacement of a hydrogen atom on a double bond, or polymerization. It has also been suggested that the appearance of an additional band at 5.72 μm, due to C==O stretching, indicates the formation of aldehydes, ketones or acids. Furthermore, changes in the absorption bands in the 10- to 11-μm region indicates cis,trans isomerization and probably formation of conjugated bonds. Determination of oxidative deterioration of lipids using IR method is simple, rapid, and requires small amounts of sample (20 mg).

van de Voort et al. [71] and Sedman et al. [72] have investigated the feasibility of employing Fourier transform infrared (FTIR) spectroscopy to assess the oxidative status or forecast the oxidative stability of oils. These authors constructed a spectral library by recording the FTIR spectra of oils spiked with various compounds representative of common oil oxidation products. Table 1 shows that each of the various types of oxidation products gives rise to discernible and characteristic absorptions in the FTIR spectrum. Similar absorption bands were detected in the spectra of oils oxidized under accelerated conditions and monitored in real time by FTIR spectroscopy. On the basis of the results of this study, the authors proposed a quantitative approach whereby the oxidative status of an oil could be determined through calibrations developed with oils spiked with appropriate compounds representative of the functional groups associated with typical oxidative end products. These concepts were subsequently put into practice with the development of a calibration for the determination of peroxide value. A similar approach may be used to develop a parallel method for evaluating p-anisidine values.

C. Chemiluminescence Spectroscopy

Burkow et al. [73] reported that hypochlorite-activated chemiluminescence could provide a useful means for evaluation of antioxidants in edible oils. Due to high sensitivity and ability to detect small changes in the degree of oxidation of lipids, this method may be employed to evaluate the effects of antioxidants on oils during low temperature storage (about 35°C) within a 24-hour period. Chemiluminescence
Table 1  Peak Positions of the Functional Group Absorptions of Reference Compounds Representative of Products Formed in Oxidized Oils

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Vibration</th>
<th>Frequency (cm(^{-1})) at peak maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>(\nu\text{OH})</td>
<td>3650 and 3550</td>
</tr>
<tr>
<td></td>
<td>(\delta\text{HOH})</td>
<td>1625</td>
</tr>
<tr>
<td>Hexanol</td>
<td>(\nu\text{ROH})</td>
<td>3569</td>
</tr>
<tr>
<td>tert-Butyl hydroperoxide</td>
<td>(\nu\text{ROOH})</td>
<td>3447</td>
</tr>
<tr>
<td>Hexanal</td>
<td>(\nu\text{RHC}==\text{O})</td>
<td>2810 and 2712</td>
</tr>
<tr>
<td></td>
<td>(\nu\text{RHC}==\text{O})</td>
<td>1727</td>
</tr>
<tr>
<td>2-Hexenal(^a)</td>
<td>(\nu\text{RHC}==\text{O})</td>
<td>2805 and 2725</td>
</tr>
<tr>
<td></td>
<td>(\nu\text{RHC}==\text{O})</td>
<td>1697</td>
</tr>
<tr>
<td></td>
<td>(\nu\text{RC}==\text{CH}==\text{HC}==\text{O})</td>
<td>1640</td>
</tr>
<tr>
<td></td>
<td>(\delta\text{RC}==\text{CH}==\text{HC}==\text{O})</td>
<td>974</td>
</tr>
<tr>
<td>2,4-Decadienal(^a)</td>
<td>(\nu\text{RHC}==\text{O})</td>
<td>2805 and 2734</td>
</tr>
<tr>
<td></td>
<td>(\nu\text{RHC}==\text{O})</td>
<td>1689</td>
</tr>
<tr>
<td></td>
<td>(\nu\text{RC}==\text{CH}==\text{HC}==\text{O})</td>
<td>1642</td>
</tr>
<tr>
<td></td>
<td>(\delta\text{RC}==\text{CH}==\text{HC}==\text{O})</td>
<td>987</td>
</tr>
<tr>
<td>4-Hexen-3-one(^a)</td>
<td>(\nu\text{RC}(==\text{O})\text{HC}==\text{CHR})</td>
<td>1703 and 1679</td>
</tr>
<tr>
<td></td>
<td>(\nu\text{RC}(==\text{O})\text{HC}==\text{CHR})</td>
<td>1635</td>
</tr>
<tr>
<td></td>
<td>(\delta\text{RC}(==\text{O})\text{HC}==\text{CHR})</td>
<td>972</td>
</tr>
<tr>
<td>Oleic acid</td>
<td>(\nu\text{RCOOH})</td>
<td>3310</td>
</tr>
<tr>
<td></td>
<td>(\nu\text{RC}(==\text{O})\text{OH})</td>
<td>1711</td>
</tr>
</tbody>
</table>

\(^a\)All double bonds in the trans form.

generally originates from electronically excited stages, such as singlet molecular oxygen in lipid peroxidation [74]. The chemiluminescence method has been tested for estimating the degree of deterioration of edible oils containing antioxidants [75] as well as for shelflife dating of fish samples [76].

D. NMR Spectroscopy

High-resolution nuclear magnetic resonance (NMR) spectroscopy makes it possible to determine various types of hydrogen atoms (protons, \(^1\text{H}\)) in triacylglycerol (TAG) molecules. This is due to the fact that hydrogen atoms in a strong magnetic field absorb energy, in the radiofrequency range, depending on their molecular environment. During oxidation of food lipids, changes occur in the environment in which protons in an oxidizing TAG molecule are located. These changes may be monitored by employing \(^1\text{H}\) NMR spectroscopy [77–80]. For this purpose, the oil is dissolved in CDCl\(_3\) and its NMR spectrum recorded (Fig. 10). The sharp signal at the extreme right side of the spectrum (high applied field) is due to tetramethylsilane (TMS) added to the solution to serve as an internal standard. The spectrum shows eight groups of signals labeled \(a\–h\). These signals are assigned: \(a\), hydrogens directly attached to double-bonded carbons (olefinic protons) and the methine proton in the glyceryl moiety (\(\delta\) 5.1–5.4); \(b\), hydrogens in the two methylene groups in the glyceryl moiety (\(\delta\) 4.0–4.4); \(c\), hydrogens in the CH\(_2\) groups attached to two double-bonded carbon atoms (diallylmethylene protons) (\(==\text{HC}==\text{CH}==\>; \delta\) 2.6–2.9);
Figure 10  $^1$H Nuclear magnetic resonance spectrum of oxidized canola oil (peak at $\delta$ 0.00 and 7.26 are for tetramethylsilane (TMS) and CHCl$_3$ protons as impurities in CDCl$_3$, respectively).

$d$, hydrogens in the three CH$_2$ groups alpha to the carboxyl groups (CH$_3$; $\delta$ 2.2–2.4); $e$, hydrogens in the CH$_2$ groups attached to saturated carbons and double-bonded carbon atoms (—CH$_2$—C—; $\delta$ 1.8–2.2); $f$, hydrogens in the CH$_2$ groups attached to the saturated carbon atoms (—C—CH$_2$—C—; $\delta$ 1.45–1.8); $g$, hydrogens in the CH$_2$ groups bonded to two saturated carbon atoms ([CH$_2$]$_2$; $\delta$ 1.1–1.45); and $h$, hydrogens in the three terminal CH$_3$ groups ($\delta$ 0.7–1.0). The relative number of protons in each group is calculated based on the integration of methylene protons of the glyceryl moiety ($\delta$ 4.0–4.4) of the TAG (four protons in the two methylene groups of the TAG moiety) molecules. The area per proton is obtained as:

\[
\text{Area per each proton} = \text{area of } b\text{-type protons}/4
\]

Since area per proton is known, one may calculate the number of protons belonging to each and every individual signal by dividing the integration number of individual signals by the area per proton. As an example, the total number of diallylmethylene protons equals the area of $c$-type protons/(the area of $b$-type protons/4).

The total number of aliphatic, olefinic, and diallylmethylene protons are calculated, from which ratios of aliphatic to olefinic protons ($R_{ao}$) and aliphatic to diallylmethylene ($R_{ad}$) protons may be obtained. These ratios increase steadily during the storage and oxidation of oils. Shahidi [79] and Wanasundara and Shahidi [80] have
shown that the ratio of olefinic to aliphatic protons, measured by NMR, decreases continuously as long as the oxidation reaction proceeds. They suggested that the NMR technique could be useful for measuring oxidative deterioration of oils containing PUFAs, even at stages beyond the point at which peroxide value profile reaches a maximum. Saito and Udagawa [78] have used this method to evaluate oxidative deterioration of brown fish meal and suggested that NMR methodology is suitable for comparing the storage conditions of the fish meal as well as estimating the effect of antioxidants in both fish meal and fish oil. These authors reported good correlations between peroxide values and NMR data. However, Wanasundara and Shahidi [80] found that linear relationships between peroxide values and NMR data were not as suitable as those of TOTOX values and NMR data. It is obvious that TOTOX values correlate better with R∞ and R6 than peroxide values since both TOTOX and NMR data estimate overall changes that occur in fatty acid profiles as reflected in both primary and secondary oxidation products of lipids. Thus, NMR methodology offers a rapid, nondestructive, and reliable technique for estimating the oxidative state of edible oils during processing and storage.

E. Chromatographic Techniques

Different chromatographic techniques have been developed and applied to quantitate oxidation products in a variety of substances, including model compounds, oils, and food lipids, subjected to oxidation under very different conditions, from room to frying temperatures. For quantification of free MA, reversed phase HPLC using ion-pairing reagent or size exclusion separation followed by monitoring the absorbance at 267 nm has been described [81–83]. MA and 4-hydroxynonenal (4-HNE) can also be derivatized with 2,4-dinitrophenylhydrazine at room temperature to form dinitrophenylhydrazone (DNP) derivatives. The DNP derivatives could be solubilized in organic solvents and separated on a reversed phase HPLC and detected at 300–330 nm, depending on the type of hydrozone formed [84–88]. GC methods to quantify MA have also been reported. The advantages of the GC methods are increased sensitivity, particularly when used with MS detection, and the possibility for simultaneous analysis of several aldehydes. Reduction of MA to 1,3-propanediol with borane trimethylamine [89] forms a butyldimethylsilyl ether, which can be analyzed by GC-MS with an HP-5 capillary column (25 m long), temperature programmed from 115°C to 165°C and [2H₆]propanediol as the internal standard. MA can also be converted to 1-methylpyrazole by reaction with N-methylhydrazine at room temperature for a 1-hour period; this derivative could be recovered by extraction with dichloromethane and analyzed on a DB-Wax capillary column (30 m long), temperature programmed from 30°C to 200°C, and a nitrogen-phosphorus detector [90,91]. Oils heated at high temperatures (oil used for frying) and (TAGs) oxidize and form polymeric TAG and hydrolytic products (e.g., diacylglycerols and fatty acids). Solid phase extraction with silica could be used to separate polar and nonpolar fractions of oxidized oils. The polar fraction can be analyzed by high-performance size exclusion column (highly cross-linked styrene–divinylbenzene copolymer) using a refractive index detector. Polar compounds are separated as an inverse order of molecular weight; TAG polymers, TAG dimers, oxidized TAG monomers, diacylglycerols, monostearine, and fatty acids [92]. It has been observed that oxidized monomers show a progressive increase during early stages of oxidation. According
to Marquez-Ruiz et al. [93], who used trilinolein (LLL) as the model compound, during early stages of oxidation LLL-oxidized monomers increase paralleled that of peroxide value, as primarily hydroperoxides were formed. The peroxides which are labile products readily degrade to a multitude of secondary products, such as oxygenated side products of the same chain length as the parent hydroperoxides. The oxidized TAGs comprise those monomeric TAGs containing at least one oxidized fatty acyl group (e.g., a peroxide group or any other oxygenated function, such as epoxy, keto, hydroxy). Therefore, determination of oxidized TAG monomers may provide a measure of both primary and secondary products of lipid oxidation.

Oxidized fatty acid methyl esters (FAMEs) could also be analyzed with a combination of silica column chromatography and high-performance size exclusion separation. The combined chromatographic analysis permits quantitation of groups of compounds (nonpolar fatty acid monomers, dimers, oxidized fatty acid monomers, and fatty acid polymers) differing in polarity of molecular weight [94]. The high-performance size exclusion chromatography–separated fractions of these oxidized fatty acids could be further analyzed on GC-MS for detection of their structural identities. The fraction of oxidized fatty acid monomers includes epoxides, ketones, and hydroperoxides as well as polyoxygenated monomeric compounds. Marquez-Ruiz and Dobarganes [94] also described that GC-MS coupled with DB-wax column and AEI-MS was useful in identifying short chain aldehydes resulting from the breakdown of lipid hydroperoxides.

REFERENCES


I. INTRODUCTION

Lipid oxidation in foods is a serious problem, difficult to overcome often and leads to loss of shelf life, palatability, functionality, and nutritional quality. Loss of palatability is due to the generation of off-flavors that arise primarily from the breakdown of unsaturated fatty acids during autoxidation. The high reactivity of the carbon double bonds in unsaturated fatty acids makes these substances primary targets for free radical reactions. Autoxidation is the oxidative deterioration of unsaturated fatty acids via an autocatalytic process consisting of a free radical chain mechanism [1]. The chain of reaction includes initiation, propagation, and termination. Propagation reactions are primarily responsible for the autocatalytic nature of autoxidation.

Autoxidation must be induced by preformed or primary hydroperoxides. The source of preformed hydroperoxides is either photosynthesized oxidation or lipoxygenase catalysis. Photosynthesized oxidation (photooxidation) involves direct reaction of light-activated, singlet oxygen with unsaturated fatty acid and the subsequent formation of hydroperoxides. Lipoxygenase catalysis involves enzymatic oxidation of unsaturated fatty acids to their corresponding hydroperoxides. Formation of primary hydroperoxides is catalyzed by either light, metals, singlet oxygen and sensitizers, or preformed hydroperoxide decomposition products.
II. ANTIOXIDANTS

In foods containing lipids, antioxidants delay the onset of oxidation or slow the rate at which it proceeds. These substances can occur as natural constituents of foods, but they also can be intentionally added to products or formed during processing. Their role is not to enhance or improve the quality of foods, but they do maintain food quality and extend shelf life. Antioxidants for use in food processing must be inexpensive, nontoxic, effective at low concentrations, stable, and capable of surviving processing (carry-through effect); color, flavor, and odor must be minimal. The choice of which antioxidant to use depends on product compatibility and regulatory guidelines [2].

Antioxidants not only extend product shelf life but reduce raw material waste, reduce nutritional losses, and widen the range of fats that can be used in specific products [3]. By extending keeping quality and increasing the number of oils that can be used in food products, antioxidants allow processors to use more available and/or less costly oils for product formulation.

A. Classification

Antioxidants can be broadly classified by mechanism of action as primary antioxidants and secondary antioxidants. Some antioxidants exhibit more than one mechanism of activity and are often referred to as multiple-function antioxidants. Chemical modes of action vary greatly, for these substances are able to function at all stages of the free radical reaction.

1. Primary Antioxidants

Primary, type 1, or chain-breaking antioxidants are free radical acceptors that delay or inhibit the initiation step or interrupt the propagation step of autoxidation. Initiation of autoxidation occurs when an \( \alpha \)-methylene hydrogen molecules is abstracted from an unsaturated lipid to form a lipid (alkyl) radical (R•) [Eq. (1)].

\[
\text{RH} \rightarrow \text{R} + \text{H}.
\]  

(1)

This highly reactive lipid radical can then react with oxygen to form a peroxy radical (ROO•) in a propagation reaction [Eq. (2)].

\[
\text{R} + \text{O}_2 \rightarrow \text{ROO} + \text{H}.
\]

(2)

During propagation, peroxy radicals react with lipid to form a hydroperoxide and a new unstable lipid radical [Eq. (3)]. This lipid radical will then react with oxygen to produce another peroxy radical, resulting in a cyclical, self-catalyzing oxidative mechanism [Eq. (4)].

\[
\text{ROO} + \text{RH} \rightarrow \text{ROOH} + \text{R}.
\]

(3)

\[
\text{R} + \text{O}_2 \rightarrow \text{ROO} + \text{H}.
\]

(4)

Hydroperoxides are unstable and can degrade to produce radicals that further accelerate propagation reactions. These reactions are typically referred to as branching steps [Eqs. (5) and (6)].

\[
\text{ROOH} \rightarrow \text{RO} + \text{OH}.
\]

(5)

\[
\text{RO} + \text{RH} \rightarrow \text{ROH} + \text{R}.
\]

(6)
Hydroperoxide degradation leads to the undesirable odors and flavors associated with rancidity in later stages of oxidation.

Primary antioxidants react with lipid and peroxy radicals and convert them to more stable, nonradical products. Primary antioxidants donate hydrogen atoms to the lipid radicals and produce lipid derivatives and antioxidant radicals (A·) that are more stable and less readily available to further promote autoxidation. As hydrogen donors, primary antioxidants have higher affinities for peroxy radicals than lipids [4]. Therefore, peroxy and oxy free radicals formed during the propagation [Eqs. (2) and (4)] and branching [Eqs. (5) and (6)] steps of autoxidation are scavenged by primary antioxidants [Eqs. (7) and (8)]. Antioxidants may also interact directly with lipid radicals [Eq. (9)].

\[
\begin{align*}
\text{ROO}^- + AH & \rightarrow \text{ROOH} + A^- \\
\text{RO}^- + AH & \rightarrow \text{ROH} + A^- \\
R^- + AH & \rightarrow RH + A^-
\end{align*}
\]

The antioxidant radical produced by hydrogen donation has a very low reactivity with lipids. This low reactivity reduces the rate of propagation, since reaction of the antioxidant radical with oxygen or lipids is very slow. The antioxidant radical is stabilized by delocalization of the unpaired electron around a phenol ring to form stable resonance hybrids. Antioxidant radicals are capable of participating in termination reactions with peroxy [Eq. (10)], oxy [Eq. (11)], and other antioxidant radicals [Eq. (12)]. The formation of antioxidant dimers (dimerization) is prominent in fats and oils and indicates that phenolic antioxidant radicals readily undergo termination reactions. This effectively stops the autocatalytic free radical chain mechanism as long as the antioxidant is present in its nonradical form.

\[
\begin{align*}
\text{ROO}^- + A^- & \rightarrow \text{ROOA} \\
\text{RO}^- + A^- & \rightarrow \text{ROA} \\
A^- + A^- & \rightarrow AA
\end{align*}
\]

Prior to initiation of autoxidation, there must be an induction period in which antioxidants are consumed and free radicals are generated. Therefore, primary antioxidants are most effective if they are added during the induction and initiation stages of oxidation when the cyclical propagation steps have not occurred. Addition of antioxidants to fats that already contain substantial amounts of peroxides will quickly result in loss of antioxidant function [5]. In addition to radical scavenging, primary antioxidants can reduce hydroperoxides to hydroxy compounds. However, the main antioxidative mechanism of primary antioxidants is radical scavenging.

Primary antioxidants are mono- or polyhydroxy phenols with various ring substitutions. Substitution with electron-donating groups ortho and para to the hydroxyl group of phenol increases the antioxidant activity of the compound by an inductive effect. These hindered phenolic antioxidants decrease the reactivity of the hydroxyl group by increasing its electron density. Substitution with butyl or ethyl groups para to the hydroxyl enhances the antioxidant activity. Because of steric hindrance, however, the presence of longer chain or branched alkyl groups at the para positions can decrease antioxidant effectiveness [6]. Substitutions of branched alkyl groups at ortho positions enhance the phenolic antioxidant’s ability to form stable resonance struc-
tures and further reduce the antioxidant radical’s ability to participate in propagation reactions.

The most commonly used primary antioxidants in foods are synthetic compounds. Examples of important primary phenolic antioxidants include butylated hydroxyanisole (BHA), butylated hydroxytoluene (BHT), propyl gallate (PG), and tertiary butylhydroquinone (TBHQ). However, a few natural components of food also act as primary antioxidants and are commonly added to foods. Tocopherols are the most commonly used natural primary antioxidants. Carotenoids are another group of natural compounds that have primary antioxidant activity, although the mechanism differs from the phenolics.

2. Secondary Antioxidants

Secondary, preventive, or type 2, antioxidants act through numerous possible mechanisms. These antioxidants slow the rate of oxidation by several different actions, but they do not convert free radicals to more stable products. Secondary antioxidants can chelate prooxidant metals and deactivate them, replenish hydrogen to primary antioxidants, decompose hydroperoxides to nonradical species, deactivate singlet oxygen, absorb ultraviolet radiation, or act as oxygen scavengers. These antioxidants are often referred to as synergists because they promote the antioxidant activity of type 1 antioxidants. Citric acid, ascorbic acid, ascorbyl palmitate, lecithin, and tarteric acid are good examples of synergists. Some of the more important types of secondary antioxidant mechanism are discussed in the following sections.

a. Chelators.

Several heavy metals with two or more valence states (Fe, Cu, Mn, Cr, Ni, V, Zn, Al) promote oxidation by acting as catalysts of free radical reactions. These redox-active transition metals transfer single electrons during changes in oxidation states.

Two mechanisms of oxidation promotion by metals have been proposed. Metals are believed to either interact with hydroperoxides or to react directly with lipid molecules. Metals are able to promote oxidation by interacting directly with unsaturated lipids [Eq. (13)] and lowering the activation energy of the initiation step of autocatalysis. However, because of thermodynamic constraints, spin barriers, and an extremely slow reaction rate, this direct interaction of metals with lipid moieties is not the main mechanism of metal catalysis [7–9].

\[ M^{n-1+} + RH \rightarrow M^{n+} + H^+ + R^\cdot \]  
(13)

Metals are known to interact with hydroperoxides and promote oxidation. Moreover, it is thought that a metal–hydroperoxide complex forms and subsequently decomposes to produce free radicals. Metals enhance the rate of decomposition of hydroperoxides and the generation of free radicals. Two metal–hydroperoxide reactions are possible.

\[ M^{n+} \text{ ROOH} \rightarrow M^{n+} + \text{H}^+ + \text{ROO}^\cdot \]  
(14)
\[ M^{n+} + \text{ROOH} \rightarrow M^{n+1+} + \text{OH}^- + \text{RO}^\cdot \]  
(15)

Reaction (15) is less significant in aqueous solution, since metals in their lower oxidation states accelerate hydroperoxide degradation more than metals in their higher oxidation states [10]. Even trace amounts of these metals promote electron transfer from lipids or hydroperoxides because the reactions [Eqs. (14) and (15)] can
be cyclical, with regeneration of the lower oxidation state of the metal. Nevertheless, reaction (15) occurs much more slowly than reaction (14) [7]. Although the metal–hydroperoxide mechanisms are generally accepted as the most important for metal catalysis of autoxidation, it is unclear whether redox-active transition metal promote lipid peroxidation directly through the formation of metal–lipid complexes or by forming peroxy and oxy radicals.

Chelation of metals by certain compounds decreases their prooxidant effect by reducing their redox potentials and stabilizing the oxidized form of the metal. Chelating compounds may also sterically hinder formation of the metal hydroperoxide complex. Citric acid (and its lipophilic, monoglyceride ester), phosphoric acid (and its polyphosphate derivatives), and ethylenediaminetetraacetic acid (EDTA) can chelate metals. EDTA forms a thermodynamically stable complex with metal ions. The metal-chelating ability of oligophosphate increases with phosphate group number up to six residues. Carboxyl groups of citric acid are thought to be responsible for binding with metals and forming complexes. Malic, tartaric, oxalic, and succinic acids bind metals in the same manner.

In addition to their antioxidant activity, many of these compounds have other unique functions as food additives. Citric acid, malic acid, and tartaric acid are important food acidulants. Phosphates are added as buffers, emulsifiers, and acidulants and water binders. Chelating antioxidants are also referred to as synergists because they enhance the activity of phenolic antioxidants. This synergism is sometimes referred to as acid synergism when the chelator is citric or other acids.

b. Oxygen Scavengers and Reducing Agents. Ascorbic acid, ascorbyl palmitate, erythorbic acid, sodium erythorbate, and sulfites prevent oxidation by scavenging oxygen and acting as reductants. Oxygen scavenging is useful in products with head space or dissolved oxygen. Reducing agents function by donating hydrogen atoms. Ascorbic acid and sulfites react directly with oxygen and eliminate it from the food product [11]. L-Ascorbic acid has strong reducing properties, and its most significant chemical property is its ability to oxidize through one- or two-electron transfers [12,13]. One-electron reactions involve an L-ascorbic acid radical (semidehydroascorbic acid). A proton is lost, whereupon a bicyclic radical, which is the intermediate leading to dehydroascorbic acid, is formed. A two-electron transfer occurs when transition metals catalyze ascorbate autoxidation. In this process, L-ascorbate and oxygen form a ternary complex with the metal catalyst. Two π electrons from L-ascorbate shift to oxygen through the transition metal. Oxidation of ascorbic acid has been reviewed by Liao and Seib [14].

Ascorbic acid represents a truly multifunctional antioxidant. Schuler [15] identified the following classes of reactions as significant to the antioxidant action of ascorbic acid in food systems: quenching of singlet oxygen, reductions of free radicals and primary antioxidant radicals, and removal of molecular oxygen in the presence of metal ions.

c. Singlet Oxygen Quenchers. Singlet oxygen is a high energy molecule that is responsible for photooxidation of unsaturated fats and the subsequent generation of hydroperoxides. Singlet oxygen quenchers deplete singlet oxygen of its excess energy and dissipate the energy in the form of heat. Carotenoids, including β-carotene, lycopene, and lutein, are active singlet oxygen quenchers at low oxygen partial pressure. Figure 1 gives an overview of lipid oxidation and the interaction of antioxidants.
Consumers are concerned about the safety of their food and about potential effects of synthetic additives on their health. Despite the superior efficacy, low cost, and high stability of synthetic antioxidants in foods, the suspicion that these compounds may act to promote carcinogenicity has led to a decrease in their use [16]. A trend toward the use of “natural” food additives in the food industry has been apparent for quite some time—a result of consumer demand. Some natural preservatives exist inherently in foods; others can be added to the product or can arise as a result of processing or cooking. Natural food antioxidants such as citric acid and ascorbic acid are used widely in the food industry. Recent research has focused on isolation and identification of effective antioxidants of natural origin.
A. Synthetic Antioxidants

Synthetic antioxidants are intentionally added to foods to inhibit lipid oxidation. Synthetic antioxidants approved for use in food include BHA, BHT, PG (also octyl and dodecyl gallate), ethoxyquin, ascorbyl palmitate, and TBHQ. The synthesis of novel antioxidants for food use is limited by rising costs of research and development, costs associated with safety assessment, and the time required to obtain regulatory approval of additives [17]. These restrictions, as well as growing consumer preference for natural food additives, has led industry to emphasize natural materials as a source of novel antioxidants.

Phenolic compounds represent some of the oldest and most frequently used antioxidants in foods. The differences in antioxidant activity of the phenolic antioxidants are due to variations in structure that directly influence physical properties. Phenols in which the aromatic ring contains alkyl groups (hindered phenols) are extremely effective antioxidants. Hindered phenols are also effective antimicrobials in foods. A good product’s characteristics ultimately determine the selection of the phenolic antioxidant. BHA and BHT are fairly heat stable and are used in heat-processed foods. PG decomposes at 148°C and is inappropriate for high temperature processing. Therefore, heat-stable TBHQ is useful in frying applications. BHA and BHT are strongly lipophilic and are used extensively in oil-in-water emulsions. BHA and BHT are also typically used together in mixtures, acting synergistically. A summary of the physical properties and applications of phenolic antioxidants is provided in Table 1.

Titles 9 and 21 of the U.S. Code of Federal Regulations (CFR) govern the use of antioxidants in meat and poultry products and in foods, respectively. Phenolic antioxidants are effective at low concentration and are often used at levels less than 0.01% [6]. The level of phenolic antioxidants permitted in the United States varies according to the product. As specified in 21 CFR, 172.110 and 172.115, limitations for BHA and BHT alone or in combination are as follows for specific products: potato granules, 10 ppm; dehydrated potato shreds, dry breakfast cereals, potato flakes, sweet potato flakes, 50 ppm; and emulsion stabilizers for shortenings, 200 ppm. At high levels, phenolic compounds become prooxidants because of their high reactivity and participation in the initiation process. Allowable limits vary greatly depending on the food product and the antioxidant. Regulations concerning synthetic antioxidants vary greatly from country to country and complicate marketing of products internationally.

Commercial antioxidant preparations are available in solid and liquid blends. Liquid blends are convenient because the antioxidant is solubilized for each addition during processing. Solvents include vegetable oils, propylene glycol, glyceryl monoleate, ethanol, and acetylated monoglycerides [5]. Antioxidant preparations typically contain mixtures of phenolic antioxidants, a synergist, and a solvent system. Some of the most important synthetic antioxidants are discussed in the following sections. Structures are provided in Figure 2.

1. Butylated Hydroxyanisole

BHA is typically used as a 9:1 mixture of 3-BHA and 2-BHA isomers [6]. The 3-isomer shows higher antioxidant activity than the 2-isomer [5]. It is a waxy, monophenolic, white solid that is fat-soluble. It is effective at preventing oxidation of
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<th>Compound</th>
<th>Molecular weight</th>
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<th>Boiling point (°C)</th>
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<td>Butylated hydroxytoluene (BHT)</td>
<td>220.356</td>
<td>White crystals</td>
<td>265</td>
<td>70</td>
<td>Insoluble in water, glycerol, and propylene glycol. Slightly soluble in mineral oil. Soluble in fats, paraffin, glycercyl monooleate, alcohol, petroleum, ether, most organic solvents</td>
<td>Less carry-through in baked and fried products than BHA. More sterically hindered than BHA. Most effective in animal fats. Slight phenol odor. Poor carry-through properties Synergistic with other antioxidants. Decomposes at frying temperatures</td>
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<td>Propyl gallate (PG)</td>
<td>212.20</td>
<td>White crystals</td>
<td>Decomposes above 148</td>
<td>Slightly soluble in water, fats, mineral oil, glyceryl monooleate. Soluble in alcohol, glycerol, propylene glycol</td>
<td>Stability of octyl and dodecyl forms greater than propyl. Discolors in presence of metals; always used in combination with a chelator. Less soluble in fats than BHA and BHT. More effective in vegetable oils than BHA and BHT. Poor carry-through in baking, but good carry-through in frying. Synergistic with other antioxidants.</td>
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<td>Tertiary butylhydroquinone (TBHQ)</td>
<td>166.22</td>
<td>White to tan crystals</td>
<td>300</td>
<td>126.5–128.5</td>
<td>Slightly soluble in water. Moderately soluble in fats, propylene glycol, glyceryl monooleate. Soluble in alcohol</td>
<td>Excellent antioxidant in vegetable oils. Does not discolor in presence of metals. Little odor Poor carry-through in baking and frying.</td>
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<td>Ethoxyquin</td>
<td>217.31</td>
<td>Yellow liquid</td>
<td>123–125</td>
<td>Soluble in most organic solvents</td>
<td>Effective in pigment retention</td>
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animal fats, but ineffective for vegetable fats. BHA has good carry-through in baking but volatilizes during frying. It is commonly added to packaging materials.

2. Butylated Hydroxytoluene

BHT is a widely used monophenolic antioxidant. This fat-soluble white, crystalline solid is appropriate for high temperature processing but is not as stable as BHA. It will volatilize and has less carry-through. BHA and BHT act synergistically to provide greater antioxidant activity than either antioxidant alone. Therefore, foods typically contain BHA/BHT mixtures at levels up to 0.02%. The postulated synergistic mechanism of BHA and BHT involves the interactions of BHA with peroxy radicals to produce a BHA phenoxy radical. The BHA phenoxy radical is then believed to abstract a hydrogen from the hydroxyl group of BHT. BHT effectively acts as a hydrogen replenisher of BHA, allowing BHA to regenerate its effectiveness. The BHT radical can then react with a peroxy radical and act as a chain terminator [7].

3. Tertiary Butylhydroxyquinone

TBHQ is a beige diphenolic powder. It is used in frying applications with highly unsaturated vegetable oils, where it has good carry-through. It is generally considered to be more effective in vegetable oils than BHA or BHT [5]. Citric acid and TBHQ show excellent synergism in vegetable oils. TBHQ is not permitted for use in foods in Canada or the European Economic Community. In the United States, TBHQ is not permitted to be combined with PG.
4. 6-Ethoxy-1,2-dihydro-2,2,4-trimethylquinoline (Ethoxyquin)

Ethoxyquin may be used as an antioxidant for the preservation of chili powder, paprika, and ground chili at levels not to exceed 100 ppm (CFR 21, 172.140). It can be used in animal products at much lower levels. A primary role of ethoxyquin in the feed industry is the protection of carotenoids. Similarly, it stabilizes the color in paprika and chili powder. In oils, it exists primarily in the form of a radical that acts as a free radical terminator. Dimerization of this radical occurs in oil and will inactivate the antioxidant.

5. Gallates

Propyl gallate (PG), octyl gallate, and dodecyl gallate are approved for use as antioxidants in foods. PG, octyl gallate, and dodecyl gallate are, respectively, the \(n\)-propyl, \(n\)-octyl, and \(n\)-dodecyl esters of 3,4,5-trihydroxybenzoic acid. Commercially, PG is the only gallate used in substantial quantity. A slightly water-soluble, white crystalline power, PG is used widely in foods for which lipid-soluble BHA, BHT, and TBHQ are not suitable. PG is not stable at high temperatures, degrading at 148°C, and it is not suitable for frying applications. Octyl and dodecyl gallate are more lipid soluble, more heat stable, and have better carry-through. Gallates are sold as mixtures with metal chelators because they will form undesirable, dark-colored complexes with iron and copper [3]. Gallates act synergistically with primary antioxidants and some secondary antioxidants, and are often included in mixed antioxidant preparations.

6. Tocopherols

\(\alpha\), \(\beta\), \(\gamma\), \(\delta\)-Tocopherols and the corresponding tocotrienols (vitamin E homologs) are natural, monophenolic antioxidant constituents of vegetable oils. Tocopherols are described in detail in Section III.B. (Natural Antioxidants).

7. Erythorbic Acid and Ascorbyl Palmitate

Erythorbic acid (\(d\)-ascorbic acid) is often used as an antioxidant in fruits and as a curing accelerator in cured meats [2]. It is very soluble in water and insoluble in oil. It has Generally Recognized as Safe (GRAS) status with the U.S. Food and Drug Administration, but unlike \(L\)-ascorbic acid, it is not a natural constituent of foods. It has minimal vitamin C activity.

Ascorbyl palmitate and ascorbyl stearate are synthetic derivatives of ascorbic acid. Ascorbyl palmitate is used in fat-containing foods because its solubility in hydrophobic media is superior to that of ascorbic acid and its salts; it is still fairly insoluble, however, and requires the aid of solubilizing magnets and/or high temperatures for solubilization [3]. Ascorbyl palmitate is usually used in combination with tocopherols. Ascorbyl palmitate has GRAS status, and the United States imposes no restrictions on usage levels. Ascorbyl palmitate is hydrolyzed by the digestive system to provide nutritionally available ascorbic acid and palmitic acid, but health claims cannot be made for its vitamin C contribution [13].

B. Natural Antioxidants

Extensive research has been dedicated to identification of antioxidants from various natural sources. Ascorbic acid and tocopherols are the most important commercial
natural antioxidants. Other sources of natural antioxidants include carotenoids, flavonoids, amino acids, proteins, protein hydrolysates, Maillard reaction products, phospholipids, and sterols. Numerous naturally occurring phenolic antioxidants have been identified in plant sources and vegetable extracts. Enzymes also play important roles as antioxidants. Processing of foods can induce the formation of antioxidants. Maillard reaction products, protein hydrolysates, fermentation products, and nitrosyl compounds from curing have been reported to possess antioxidant activity. Natural antioxidants allow food processors to produce stable products with “clean” labels that tout all-natural ingredients. However, these products can have several drawbacks, including high usage levels, undesirable flavor and/or color contributions, and lack of stability due to low antioxidant efficiency.

The safety of natural antioxidants should not be taken for granted. Cautions must be heeded, since numerous natural products are potential carcinogens, mutagens, or teratogens, and the safety of many natural compounds with antioxidant activity has not been established. A case in point is nordihydroguaiaretic acid (NDGA), which was used extensively as an antioxidant earlier in this century. NDGA is a natural constituent of the creosote bush, which was removed from GRAS status when unfavorable toxicological results were reported. Regardless of the politics surrounding the issue of safety and “natural” additives, natural antioxidant products are commercially important and desired by the consumer. The main advantage of substances naturally present in foods is that the burden of proof of safety may be less rigorous than that required for synthetic products. No safety testing is required if the antioxidant is a natural constituent of GRAS ingredients. In addition, some natural antioxidants derived from spices, herbs, and Maillard reactions can be listed as flavorants rather than antioxidants, a technical distinction that serves to exempt the substances from safety testing requirements.

1. Tocopherols and Tocotrienols

Tocopherols and tocotrienols (Fig. 3) comprise the group of chromanol homologs that possess vitamin E activity in the diet. They are natural monophenolic compounds with varying antioxidant activities. Eight naturally occurring homologs are included in the vitamin E family [12]. They are fat-soluble 6-hydroxychroman compounds. The α-, β-, γ-, and δ-tocopherols are characterized by a saturated side chain consisting of three isoprenoid units. The corresponding tocotrienols (α, β-, γ-, and δ-) have double bonds at the 3′, 7′, and 11′ positions of the isoprenoid side chain. Only RRR isomers are found naturally. Synthetic α-tocopherol (all-rac-α-tocopherol) consists of eight stereoisomers found in equal amounts in the synthetic mixture. Biologically, RRR-α-tocopherol is the most active vitamin E homolog. Tocopherols and tocotrienols are widely distributed in the plant kingdom, with vegetable oils providing the most concentrated source of vitamin E. Tocotrienols are less common but are present in palm oil, rice bran oil, cereals, and legumes. Palm oil has a unique vitamin E profile, providing tocotrienols in higher concentrations than other food sources. Tocopherols and tocotrienols are retained throughout the edible oil refining process, although there is some loss during the deodorization step. Worldwide, the main commercial source of natural tocopherols is in the soybean oil refining industry. Commercial natural antioxidant preparations prepared from soybean oil typically consist of greater than 80% γ- and δ-tocopherol [2]. Synthetic tocopherols are commercially available and vary in isomeric form. Tocopherols provide a useful natural
antioxidant source for foods marketed under an “all-natural” label. They are permitted in foods according to GMP regulations (21 CFR 182.3890). Natural tocopherols are limited to 0.03% (300 pp in animal fats) (9 CFR 318.7). Because of the high vitamin E content of most vegetable oils, addition of tocopherols can lead to prooxidant activity [18].

The antioxidative mechanism of \(\alpha\)-tocopherol is well understood. \(\alpha\)-Tocopherol donates a hydrogen to a peroxy radical resulting in a \(\alpha\)-tocopheryl semiquinone radical [Eq. (16)]. This radical may further donate another hydrogen to produce methyltocopherylquinone [Eq. (17)] or react with another \(\alpha\)-tocopheryl semiquinone radical to produce an \(\alpha\)-tocopherol dimer [Eq. (18)]. Higher polymeric forms can then form.

\[
\text{ROO}^\cdot + \alpha\text{-tocopherol} \rightarrow \text{ROOH} + \alpha\text{-tocopheryl semiquinone}^\cdot \quad (16)
\]

\[
\alpha\text{-tocopheryl semiquinone}^\cdot + \text{ROO}^\cdot \rightarrow \text{ROOH} + \text{methyltocopherylquinone} \quad (17)
\]

\[
\alpha\text{-tocopheryl semiquinone}^\cdot + \alpha\text{-tocopheryl semiquinone}^\cdot \rightarrow \alpha\text{-tocopherol dimer} \quad (18)
\]

The methyltocopherylquinone is unstable and will yield \(\alpha\)-tocopherylquinone. The \(\alpha\)-tocopherol dimer continues to possess antioxidant activity. Numerous other
decomposition products with various degrees of antioxidant activity can arise from 
oxidation of tocopherols [15]. The antioxidant activity of the tocols and tocotrienols 
increases from $\alpha$ through $\delta$. $\alpha$-Tocopherol and $\alpha$-tocotrienol are fully substituted 
benzoquinone derivatives and are the most effective antioxidants. $\alpha$-Tocopherol is 
highly reactive with peroxy radicals and prevents them from participating in prop-
agation reactions. The $\alpha$-tocopheroxyl radical is stable because a resonance structure 
forms on the benzoquinone ring. A comprehensive review of the fundamental anti-
oxidant chemistry of tocopherols and tocotrienols has been compiled by Kamal-Eldin 
and Appelqvist [19].

2. Ascorbic Acid and Ascorbate Salts

L-Ascorbic acid, or vitamin C, is ubiquitous (Fig. 4) in nature as a component of 
plant tissues and is produced synthetically in large quantities. Ascorbic acid is at-
tractive as an antioxidant because it has GRAS status with no usage limits, is a 
natural or nature-identical product, and is highly recognized as an antioxidant nutrient 
by the consumer. In some food products it is also a flavorant and acidulant. However, 
in foods that are heat-treated, ascorbic acid can participate in nonenzymatic browning 
and may be degraded through reductone reactions.

Ascorbic acid acts as a primary or a secondary antioxidant. In vivo, ascorbic 
acid donates hydrogen atoms as a primary antioxidant. Ascorbic acid is also capable 
of scavenging radicals directly by converting hydroperoxides into stable products. 
Ascorbic acid is an important antioxidant in plant tissue and is essential for the 
prevention of oxidative cellular damage by hydrogen peroxide [20]. In foods, ascor-
bic acid is a secondary antioxidant with multiple functions. Ascorbic acid can scaven-
ge oxygen, shift the redox potential of food systems to the reducing range, act 
synergistically with chelators, and regenerate primary antioxidants [21]. The vitamin 
is commonly used as a synergist to donate hydrogen to primary antioxidants such 
as tocopherol. Tocopheroxyl radicals are reduced back to tocopherol by ascorbic acid 
[Eq. (19)].

$$\text{Tocopheroxyl radical} + \text{ascorbic acid} \rightarrow \text{tocopherol} + \text{dehydroascorbic acid}$$

(19)

Ascorbic acid oxidizes through one- or two-electron transfers that are due to its 
enediol structure [12,13]. It is a reductone and has a high affinity for oxygen. The 
2- and 3-positions of ascorbic acid are unsubstituted. Oxidation of ascorbic acid 
occurs in two steps, with monohydroperoxide as an intermediate to the formation of 
dehydroascorbic acid.

Ascorbic acid and its salts (sodium ascorbate and calcium ascorbate) are water-
soluble and are not applicable as antioxidants for oils and fats. They are used exten-
sively to stabilize beverages. In the United States, ascorbyl palmitate is used in fat-
containing foods because its lipid solubility is superior to that of ascorbic acid. 
Ascorbyl palmitate has GRAS status, and there are no restrictions on usage levels.

3. Carotenoids

Carotenoids (Fig. 4) are yellow, orange, and red lipid-soluble pigments that are ubiq-
uitous in green plants and in fruits and vegetables. They are 40-carbon isoprenoids 
or tetraterpenes with varying structural characteristics. The two classes of carotenoids 
are carotenes and xanthophylls. Carotenes (e.g., $\beta$-carotene and lycopene) are poly-

Figure 4  Structures of ascorbic acid and related compounds and carotenoids.

eine hydrocarbons that vary in degree of unsaturation. Xanthophylls (e.g., astaxanthin and canthaxanthin) are synthesized from carotenes by hydroxylation and epoxidation reactions and therefore contain oxygen groups [22]. About 10% of the 600 or so identified carotenoids have the biological activity of vitamin A and are referred to as provitamin A compounds. β-Carotene is the most abundant of the provitamin A carotenoids found in food and also has the highest vitamin A activity [23]. β-Car-
otene is a polyene, synthesized from eight isoprene units. It has an intense orange-red color and, as a food additive, is used primarily as a colorant for oils and fats. Carotenoids can act as primary antioxidants by trapping free radicals or as secondary antioxidants by quenching singlet oxygen. Carotenoids are typically secondary antioxidants in foods. However, in the absence of singlet oxygen (low oxygen partial pressure), carotenoids may also prevent oxidation by trapping free radicals and acting as chain-breaking antioxidants.

Singlet oxygen is unstable and can react with lipids to produce free radicals. In the presence of \( \beta \)-carotene, singlet oxygen will preferentially transfer energy to \( \beta \)-carotene to produce triplet state \( \beta \)-carotene [Eq. (20)]. The transfer of energy from singlet oxygen to carotenoid takes place through an exchange electron transfer mechanism [22].

Triplet state \( \beta \)-carotene releases energy in the form of heat, and the carotenoid is returned to its normal energy state [Eq. (21)]. Carotenoids are very effective quenchers: one carotenoid molecule is able to interact with numerous singlet oxygens. \( \beta \)-Carotene, for example, can quench up to 1000 molecules of singlet oxygen [23].

\[
\begin{align*}
1^\text{O}_2 + \beta \text{-carotene} & \rightarrow 3^\text{\beta \text{-carotene*}} + 3^\text{O}_2 \\
3^\text{\beta \text{-Carotene*}} & \rightarrow \beta \text{-carotene} + \text{heat}
\end{align*}
\]  

(20) (21)

The ability of carotenoids to quench singlet oxygen is directly related to the number of carbon double bonds in the compound. Carotenoids with nine or more conjugated double bonds are very effective antioxidants. \( \beta \)-Carotene, isozeaxanthin, and lutein are all effective singlet oxygen quenchers. Xanthophylls are not as efficient at scavenging singlet oxygen because of the addition of functional groups to the hydrocarbon structure.

Carotenoids are known to scavenge free radicals at low oxygen pressures (<150 mm Hg) and to act as primary antioxidants in vitro [10,21]. The conjugated double bonds of carotenoids are very susceptible to attack by peroxy radicals. \( \beta \)-Carotene is capable of reacting with peroxy radicals to produce a resonance-stabilized carotene product [Eq. (22)]. The unsaturated structure of carotene allows delocalization of electrons in the radical. This carotene radical can then participate in termination reactions [Eq. (23)] and divert damaging peroxy radicals to less deleterious side reactions [23].

\[
\begin{align*}
\text{Carotene} + \text{ROO}^- & \rightarrow \text{carotene}^- \\
\text{carotene}^- + \text{ROO}^- & \rightarrow \text{termination product}
\end{align*}
\]  

(22) (23)

Antioxidant activity of carotenoids has been the subject of much research. Lutein, lycopene, and \( \beta \)-carotene have been reported to inhibit photooxidation of purified oils [24,25]. Anatto color, containing the carotenoid bixin, has been shown to have antioxidant activity [26]. Combinations of carotenoids and tocopherols act synergistically [26,27]. Carotenoids are very unstable, and care must be taken when they are added to processed foods. \( \beta \)-Carotene is difficult to apply as an antioxidant because it is not easily soluble in most common solvents and is very highly reactive [13]. Carotenoid stability is affected by oxygen, heat, pH, light, and the presence of metals. With improved stability, carotenoids could see widespread use as antioxidants.
4. Enzymatic Antioxidants

Glucose oxidase, superoxide dismutase, catalase, and glutathione peroxidase act as antioxidants by removing from the lipid environment either oxygen or highly oxidative species. The enzymes just named act biologically to eliminate cellular free radicals, to keep reactive oxygen species at low concentrations, and to catalyze the destruction of hydrogen peroxide. Thus, they constitute an important biological defense mechanism against free radical damage. Glucose oxidase is an enzyme that removes oxygen by using it to produce gluconic acid and hydrogen peroxide from glucose. Commercial glucose oxidase systems include catalase to hydrolyze the hydrogen peroxide [6].

Superoxide dismutase removes superoxide radicals (O$_2^-$) by converting them to triplet oxygen (O$_2^3$) [Eq. (24)]:

$$\text{O}_2^- + 2\text{H}^+ \rightarrow \text{H}_2\text{O}_2 + ^3\text{O}_2$$  \hspace{1cm} (24)

Catalase then converts the hydrogen peroxide to water [Eq. (25)]:

$$\text{H}_2\text{O}_2 \rightarrow \text{2H}_2\text{O} + ^3\text{O}_2$$  \hspace{1cm} (25)

Glutathione dehydrogenase catalyzes the oxidation of glutathione (γ-Glu-Cys-Gly, GSH) in the presence of dehydroascorbic acid. Dehydroascorbic acid acts as a hydrogen acceptor and is returned to its active, free radical scavenging form, ascorbic acid. Glutathione peroxidase oxidizes glutathione in the presence of hydroperoxide. The sulfhydryl group on one GSH will react with the sulfhydryl group of another GSH to produce oxidized glutathione (GSSG) with a disulfide bond [Eq. (26) [7].

$$\text{ROOH} + 2\text{GSH} \rightarrow \text{ROH} + \text{H}_2\text{O} + \text{GSSG}$$  \hspace{1cm} (26)

Glucose oxidase, catalase, and superoxide dismutase are used commercially as antioxidants in various foods.

5. Proteins and Related Substances

Numerous amines, amino acids, peptides, and protein hydrolyzates have antioxidant activity. A comprehensive list of all the proteinaceous antioxidant sources is not presented here. Amines have been shown to possess antioxidant activity. Recently, spermine and spermidine isolated from fish sources were used to inhibit fish oil oxidation [28]. Numerous amines such as hypoxanthine and xanthine can be readily isolated from marine sources. Amino acids have chelating abilities, but also exhibit antioxidant activity when used alone. Glycine, methionine, histidine, tryptophan, proline, and lysine are effective antioxidants in oil [6].

Proteins and protein hydrolysates possess antioxidative factors. Iron-binding proteins such as ferritin and transferritin have antioxidant function. Histidine-containing peptides such as carnosine [29], as well as synthetic peptides [30] and peptides obtained from protein hydrolysis [31,32], possess antioxidant activity. Glutathione tripeptide (γ-Gly-Cys-Gly, GSH) has antioxidant activity, which is mediated by the sulfhydryl group of cysteine and glutathione peroxidase. The sulfhydryl group on one GSH will react with the sulfhydryl group of another GSH to produce oxidized glutathione (GSSG) with a disulfide bond. Glutathione/glutathione peroxidase was effective in preventing lipid oxidation in a minced mackerel system [33]. In addition, GSH can form disulfides with proteins or other thiols [34]. It is capable of acting as
a free radical scavenger by forming a thiol radical. GSH also donates electrons for
the reduction of dehydroascorbate to ascorbic acid in the presence of glutathione
dehydrogenase. Ascorbic acid is then returned to its active, free radical scavenging
form. The enzyme has been purified from wheat flour, which has a relatively high
activity [7].

6. Maillard Reaction Products

Maillard reaction products (MRPs) are an excellent example of natural, process-
induced oxidation inhibitors that arise as a result of cooking [35]. MRPs are formed
during the cooking of low-moisture foods at temperatures above 80°C. They are
produced from the reaction of amines and reducing sugars. Lipids, vitamins, and
other food constituents also participate in Maillard reactions. MRPs are presumed to
be safe because they occur naturally as products in cooked foods. They are often
used as bases for flavorants and gravies. The brown pigment associated with MRPs
can be an advantage in cooked products where brown color is desirable. Use of
MRPs as a source of antioxidants in foods has been intensively studied, but our
understanding of the compounds responsible for the antioxidant activity is incomplete
and the mechanisms of action are unknown. Identification of the compounds re-
sponsible for antioxidant activity has proved difficult because of the complexity of
the Maillard reaction, the vast number and variety of MRPs, and the diversity of the
model systems that can be studied. MRPs have been shown to have antioxidant
activity in model systems as well as in some fat-containing foods [36–39]. Lingnert
and Eriksson [39] used processing parameters and the Maillard reaction to prevent
oxidation in cookies, and Sato et al. [40] inhibited warmed-over flavor in cooked
beef with MRPs.

Because of the conflicting views present in the literature, it is difficult to state
conclusively which of the numerous MRPs are actually responsible for antioxidant
activity. It is even more difficult to attempt to describe the mechanism of action of
these suspected antioxidants. Investigators have shown a correlation between color-
less, low molecular weight, intermediate MRPs and antioxidant activity [41–44].
Evans et al. [45] showed a correlation between antioxidant activity and reductone
levels in MRP mixtures. In contrast, Kirigaya et al. [36] showed that antioxidant
activity was proportional to the color intensity of MRP and proposed that nondi-
alyzable, high molecular weight melanoids, which inhibit hydroperoxide and car-
bonyl compound formation, were responsible for antioxidant activity. By completely
oxidizing the reductones in the MRP solution with DPI butanol, these investigators
also demonstrated that reductones contributed little antioxidant activity in their model
system [36]. Other researchers found a direct relationship between color intensity of
Maillard reaction solutions and antioxidant activity [38]. Yamaguchi et al. [38] also
showed increases in antioxidant activity with increases in melanoidin formation.

Theories on the mechanism of antioxidant activity of MRPs conflict as well.
Kawakishi et al. [46] hypothesized that the protective effects of melanoids against
autoxidation were likely to depend on their ability to chelate metals. Amadori com-
pounds may behave like reductones, which inhibit autoxidation. Eichner [44] believes
that MRP intermediates may scavenge oxygen. These conflicting reports likely reflect
the different reaction conditions used in the experiments and the multiple antioxi-
dative functions exhibited by MRPs with different mechanisms of action. Because
of the overwhelming complexity of even model systems of the Maillard reaction, it
would be imprudent to discount any of these theories as to the nature and activity of antioxidant MRPs.

7. Phospholipids

The antioxidative action of phospholipids is not well understood. It is likely that antioxidant activity differs among the various phospholipids as a result of the wide variance in functional groups and structures. Possible actions include regeneration of primary antioxidants, metal chelation, and decomposition of hydroperoxides. Phospholipids have been shown to be synergists. Moreover, phosphatidylcholine (PC), phosphatidylethanolamine (PE), and phosphatidylserine (PS), display antioxidant activity that is possibly linked to chelating ability [47]. Lecithin, once an important commercial antioxidant, now sees limited use because of inefficiency as an antioxidant and poor heat stability. Burkow et al. [48] found lecithin to have antioxidant activity in cod liver oil.

8. Sterols

Sterols have been documented to have antioxidant activity. It is thought that sterols interact with oil surfaces and inhibit oxidation. Sterols may be oxidized at oil surfaces and inhibit propagation by acting as hydrogen donors. Maestroduran and Borjapadilla [49] have reviewed sterol antioxidants and provided information about recent patents.

9. Sulfur Dioxide and Other Sulfites

Sulfites are reducing agents that are weak antioxidants in foods. Sulfites such as sulfur dioxide, sodium sulfite, and sodium, potassium, and metabisulfites are used to prevent flavor and color degradation in beverages and fruits. Sulfites react with molecular oxygen to form sulfates. They also act as reducing agents that promote the formation of phenols from quinones, thereby preventing browning reactions.

10. Gums

Polysaccharides have been studied for their antioxidant effects. Gums are primarily used for their texture-enhancing effects, but they also possess antioxidant activity, which may be due to metal chelation and oxygen consumption, and their viscosity-increasing effects [50,51]. Guaiaconic acids present in the resin of the *Gaujacum officinale* L. tree are responsible for the antioxidant activity of this food additive [15]. Guaiac gum was commonly used to preserve refined animal fats but has limited use now. It is not as potent as synthetic antioxidants, has poor heat stability, and is fairly expensive. Xanthan gum, pectin, guar gum, and tragacanth gum are recognized as antioxidants.

11. Antioxidants in Plants

Antioxidant components of plants include vitamin E homologs, carotenoids, proteins, and many other compounds. Plants produce a diverse assortment of phenolic metabolites that readily undergo oxidation and have the potential to minimize effects of autoxidation. Several phenolics in addition to vitamin E (Sec. III.B.1) have shown potential for use as food antioxidants or are already serving as such. Common plant phenolic antioxidants include gallic acid (as a constituent of polymeric gallotannins and ellagitannins) and protocatechuic acids, phenylpropanoids, and mixed-pathway metabolites such as alkyl ferulates, flavonoids, and suberins [52]. Gallic acid is typ-
Flavonoids (Fig. 5) are secondary products of plant metabolism and consist of anthocyanins, catechins, flavones, flavonols, isoflavones, and proanthocyanidins. Several of the flavonoids have antioxidant activity related to their ability to chelate metals. Polyvalent phenol structures in flavonoids can form complexes with metal ions. Flavonoids also act as primary antioxidants and superoxide anion scavengers [6]. These compounds are responsible for the antioxidant activity reported in many plant and spice extracts. Phenolic acids are structurally related to flavonoids and serve as precursors of their biosynthesis. Phenolic acids such as hydroxycinnamic (cafeic, \( p \)-coumaric, ferulic, and sinapic acids), hy-
droxycoumarin (scopoletin), and hydroxybenzoic acids (4-hydroxybenzoic, ellagic, gallic, gentisic, protocatechuic, salicylic, and vanillic acids) are phenolic compounds that can form metal complexes. Antioxidant activity of these compounds varies greatly and is also dependent on the food system. Hydroxycinnamic acid esters were found to be more active than the free acids in model systems involving linoleic acid [53].

Flavones (apigenin, chrysoeriol, diosmetin, isovitexin, luteolin, and nobiletin) and flavonols (gossypetin, isorhamnetin, kaempferol, myricetin, robinetin, and quercetin) occur in fruits as glycosides. These compounds are also prevalent in vegetables, tea, and wine [54]. Quercitin has gained attention as a very potent antioxidant [55]. It has been shown to be very effective in linoleic model systems [56]. Phenolic acids, flavonoids, and other phenolics have potential as food antioxidants. Contents of specific phenolic compounds in plants can be very low, requiring large amounts of raw material to obtain sufficient amounts of these antioxidants. Usage can be limited because the compounds are often present in the form of glycosides and are not soluble in oil. In addition, some flavonoids are toxic.

b. Other Natural Sources of Antioxidants. Numerous plants have been identified as sources of phenolic compounds with antioxidant activity. The list of natural antioxidants is growing as a result of the amount of research that is being conducted to isolate and identify these compounds in plants. A comprehensive listing of all the sources of antioxidant compounds identified in plant materials is beyond the scope of this chapter. Recent reviews by Pratt [56] and Pratt and Hudson [57] contain detailed information on numerous phenolic compounds. Some recent research findings are given below. The diversity of plant sources and compounds considered to be antioxidants is evident even from this brief description.

For example, flavonoid derivatives have been recently identified as potent antioxidants found in apples [58,59] and chrysanthemum [60]. Antioxidant activity has been correlated to flavonoids found in rice, buckwheat, barley, and malt [61–64]. Other examples of antioxidants include tannins from bark, lignans from papua mace, and capsaicin from peppers. Compounds with antioxidant activity isolated from high alpine plant species [65], marine sources [66], and wood smoke [2] have also been characterized.

Tea. Tea extracts are a source of natural antioxidants. Tea catechins (Fig. 6) have potent antioxidant activity [67–71]. Extracts of green and black tea contain epicatechin, epicatechin gallate, epigallocatechin, epigallocatechin gallate, and galallocatechins [72]. During the fermentation process required to produce black tea, catechins are oxidized to produce flavins and flavin gallates that have antioxidant activity [73]. Tea antioxidants have been patented for use in several food products [74–76].

Sesame seed compounds. Several antioxidative compounds have been isolated and identified from sesame seed oil (Fig. 7). The compounds are lignanphenols such as pinoresinol sesaminol, sesamol (and its dimer), and sesamolinol [77–79]. Sesame seed oil has a much longer shelf life than many edible oils because of the presence of these phenolic compounds [80].

Soybean. Soybeans have several antioxidative constituents that include soy proteins [81]. Soy and other vegetable hydrolysates contain phenolic compounds with antioxidant activity. Fermented soy products contain isoflavones and genistein, which
inhibit oxidation. Diets containing soy provide genestin, which is a phytoestrogen. Dietary estrogens are structurally similar to endogenous estrogens and mimic their action by binding to the estrogen receptor [82]. Dietary estrogen can produce biological effects similar to those of the endogenous estrogen. The role of dietary estrogens such as genistein in preventing or enhancing the progression of such chronic diseases as breast cancer, heart disease, or bone loss has not been clarified at this time. However, the interest in the biological effects of dietary estrogens is great both in the research community and among consumers.

Herbs and spices/spice extracts. Spices and herbs have been used as flavorants for thousands of years. The antioxidant activity of spices and herbs is thought to be primarily due to the presence of phenolic compounds and especially phenolic acids and flavonoids. The strong flavor of spices and herbs precludes their use in many food products. Researchers have tried to identify and isolate specific antioxidant components of spices and herbs that do not contribute undesirable flavor or color to foods. To date, only rosemary and sage (Perilla plants) are commercially available as flavorless, odorless, and colorless antioxidant extracts. Thyme, fenugreek [83], and turmeric [84] possess antioxidative components and may provide sources for commercial products.

Rosemary antioxidants (Fig. 8) have been used in processed foods for decades. Currently, concentrated rosemary extracts are available that do not impart flavor or
color to foods. The use of these products is increasing significantly with the rising consumer demand for natural food additives [2]. These extracts are substantially more expensive than synthetic alternatives and require higher usage levels because of lower efficiencies. The diterpene phenolics carnosol and carnosic acid have been shown to be major antioxidant constituents of rosemary extracts [85,86]. These compounds are as active as $\alpha$-tocopherol in bulk oils [87]. Carnosic acid was even more potent than BHA and BHT in soybean oil [88].

12. Nutritional Aspects

Antioxidant nutrients including vitamin C (ascorbic acid), vitamin E, carotenoids (such as $\beta$-carotene), and natural phenolic constituents of foods are being touted as free radical scavengers that may act to prevent cancer, heart disease, and cataracts [89]. Some researchers suggest that synthetic antioxidants in the diet may also prevent cancer formation. Free radicals, produced in the body as a natural product of oxidative reactions, can cause oxidation of cell lipids and DNA damage that may lead to serious diseases. Dietary antioxidants are thought to scavenge these free radicals.
radicals, prevent them from damaging cells and DNA, and possibly reduce oxidized fatty acids or mutagens that lead to heart disease or cancer. Epidemiological studies have shown trends suggesting that antioxidants may be beneficial in disease prevention. However, a cause-and-effect relationship cannot be determined in humans because lifestyle factors interfere with interpretation of research results.

REFERENCES


Antioxidant Mechanisms

ERIC A. DECKER

University of Massachusetts, Amherst, Massachusetts

I. INTRODUCTION

Krinsky [1] has defined biological antioxidants as “compounds that protect biological systems against the potentially harmful effects of processes or reactions that cause extensive oxidations.” While food lipids are derived from biological systems, the ultimate purpose of food antioxidants is different, since they are used to inhibit oxidative reactions that cause deterioration of quality (e.g., of flavor, color, nutrient composition, texture). With this goal in mind, food antioxidants can be defined as any compounds serving to inhibit oxidative processes that deteriorate the quality of food lipids. Antioxidant mechanisms that fit this definition include free radical scavenging, inactivation of peroxides and other reactive oxygen species, chelation of metals, and quenching of secondary lipid oxidation products that produce rancid odors.

Reactive oxygen species and free radicals are produced by both enzymic and nonenzymic reactions. Therefore, foods usually contain endogenous antioxidants to protect against oxidative damage. These antioxidant systems often contain several distinctively different antioxidants for protection against different prooxidative compounds, including transition metals, heme-containing proteins, enzymes, photosensitizers, and numerous sources of free radicals. Since prooxidants are both water- and lipid-soluble, endogenous antioxidant systems in foods are usually biphasic. Such multicomponent and biphasic antioxidants represent nature’s own hurdle technology antioxidant system.

This chapter covers the basic mechanisms by which antioxidants influence oxidative reactions: inactivation of free radicals, control of oxidation catalysts, inactivation of oxidation intermediates, and interactions between antioxidants and secondary lipid oxidation products.
II. INACTIVATION OF FREE RADICALS

Antioxidants can slow lipid oxidation by inactivating or scavenging free radicals, thus inhibiting initiation and propagation reactions. Free radical scavengers (FRS) or chain-breaking antioxidants are capable of accepting a radical from oxidizing lipids species such as peroxyl (LOO⋅) and alkoxyl (LO⋅) radicals by the following reactions [2]:

\[ \text{LOO} \cdot \text{or LO} \cdot + \text{FRS} \rightarrow \text{LOOH or LOPH} + \text{FRS} \cdot \]

FRS primarily react with peroxyl radicals for several reasons: (a) because propagation is a slow step in lipid oxidation, meaning that peroxyl radicals are often found in the greatest concentration of all radicals in the systems; and (b) because peroxyl radicals have lower energies than radicals such as alkoxyl radicals [3] and thus react more readily with the low energy hydrogens of FRS than with polyunsaturated fatty acids, and (c) because FRS, being generally found at low concentrations, do not compete effectively with initiating radicals (e.g., ·OH) [4]. A FRS thus inhibits lipid oxidation by more effectively competing with other compounds (especially unsaturated fatty acids) for peroxyl radicals.

Chemical properties, including hydrogen bond energies, resonance delocalization, and susceptibility to autoxidation, will influence the antioxidant effectiveness of an FRS. Initially, antioxidant efficiency is dependent on the ability of the FRS to donate hydrogen to the free radical. As the hydrogen bond energy of the FRS decreases, the transfer of the hydrogen to the free radical is more energetically favorable and thus more rapid. The ability of a FRS to donate a hydrogen to a free radical can be predicted from standard one-electron reduction potentials [3].

Any compound that has a reduction potential lower than the reduction potential of a free radical (or oxidized species) is capable of donating a hydrogen to that free radical unless the reaction is kinetically unfeasible (Table 1). For example, FRS including α-tocopherol \((E'^{\circ} = 500 \text{ mV})\), urate \((E'^{\circ} = 590 \text{ mV})\), catechol \((E'^{\circ} = 530 \text{ mV})\), and ascorbate \((E'^{\circ} = 282 \text{ mV})\) all have reduction potentials below peroxyl radicals \((E'^{\circ} = 1000 \text{ mV})\) and are therefore capable of donating a hydrogen atom to the peroxyl radical to form a peroxide. Standard reduction potentials can also be used to predict the ease with which a compound can donate its hydrogen to a radical.

<table>
<thead>
<tr>
<th>Couple</th>
<th>(E'^{\circ}) (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HO⋅, H⁺/H₂O</td>
<td>2310</td>
</tr>
<tr>
<td>RO⋅, H⁺/ROH</td>
<td>1600</td>
</tr>
<tr>
<td>ROO⋅, H⁺/ROOH</td>
<td>1000</td>
</tr>
<tr>
<td>PUFA⋅, H⁺/PUFA-H</td>
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<tr>
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<td>Catechol-O⋅, H⁺/catechol-OH</td>
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</tr>
<tr>
<td>α-Tocopheroxyl⋅, H⁺α-tocopherol</td>
<td>500</td>
</tr>
<tr>
<td>Ascorbate⋅, H⁺/ascorbate</td>
<td>282</td>
</tr>
</tbody>
</table>

*Source:* Adapted from Ref. 3.
For instance, the hydrogen of the hydroxyl group on α-tocopherol has a lower reduction potential than the methylene-interrupted hydrogen of a polyunsaturated fatty acid ($E^{\circ}\text{H}_2 = 600 \text{ mV}$), thus allowing the α-tocopherol to react with peroxyl radicals more rapidly than is possible for unsaturated fatty acids.

The efficiency of the FRS is also dependent on the energy of the resulting free radical scavenger radical (FRS·). If the FRS· is a low energy radical, then the likelihood that the FRS· will catalyze the oxidation of other molecules decreases. The most efficient FRS have low energy radicals as a result of resonance delocalization (Fig. 1) [5,6]. This can again be seen in standard reduction potentials, where FRS such as α-tocopherol and catechol have lower reduction potentials than polyunsaturated fatty acids and therefore do not efficiently abstract hydrogens from unsaturated fatty acids (Table 1) [3]. Efficient FRS also produce radicals that do not react rapidly with oxygen to form peroxides. When a radical scavenger forms peroxides during oxidation, it is likely that it will autoxidize, thus depleting the system of the free radical scavenger. Also, FRS peroxides can decompose into additional radicals species, which could further promote oxidation. Thus, formation of FRS peroxides can result in consumption of the antioxidant with no net decrease in free radicals numbers [4].

FRS radicals may undergo additional reactions that remove radicals from the system; examples include termination reactions with other FRS· or lipid radicals to form nonradical species (Fig. 2). This means that each FRS is capable of inactivating at least two free radicals, the first being inactivated when the FRS interacts with the peroxyl radicals and the second when the FRS· enters a termination reaction with another peroxyl radical.

Phenolics possess many of the properties of an efficient FRS. Hydrogen donation generally occurs through the hydroxyl group, and the radical subsequently formed is stabilized by resonance delocalization throughout the phenolic ring structure. The effectiveness of phenolic FRS can be increased by substitution groups. Alkyl groups in the ortho and para positions enhance the reactivity of the hydroxyl hydrogen toward lipid radicals; bulky groups at the ortho position increase the stability of phenoxy radicals; and a second hydroxy group at the ortho or para position stabilizes the phenoxy radical through an intramolecular hydrogen bond [5]. In foods, the efficiency of phenolic FRS depends on additional factors. Besides chemical reactivity, factors such as volatility, pH sensitivity, and polarity can influence the retention and activity of the FRS in stored and processed foods [6].

A. Tocopherols

Tocopherols are a group of phenolic FRS isomers originating in plants and eventually ending up in animal foods via the diet [7]. Interactions between tocopherols and lipid peroxyl radicals lead to the formation of a hydroperoxide and several resonance structures of tocopheroxyl radicals (Fig. 3) [6]. Tocopheroxyl radicals can interact with other compounds or with each other to form a variety of products. The types and amounts of these products depend on oxidation rates, radical species, lipid state (e.g., bulk vs. membrane lipids), and tocopherol concentration.

Under conditions of low oxidation rates in lipid membrane systems, tocopheroxyl radicals primarily convert to tocopherolquinone. Tocopherolquinone can form when the interaction of two tocopheroxyl radicals leads to the formation of to-
Figure 1  Resonance stabilization of a free radical by a phenolic. (Adapted from Ref. 5.)
Figure 2  Mechanism by which one phenolic free radical scavenger can inactivate two peroxyl radicals.

copherylquinone and the regeneration of tocopherol (Fig. 4) [6]. Formation of tocopherylquinone is also thought to occur by the transfer of an electron from a tocopheroxyl radical to a phospholipid peroxyl radical, to form a phospholipid peroxyl anion and a tocopherol cation. The tocopherol cation hydrolyzes to $8\alpha$-hydroxytocopherone, which rearranges to tocopherylquinone (Fig. 5) [8].

Under condition of more extensive oxidation, high concentrations of peroxyl radicals can favor the formation of tocopherol–peroxyl complexes. These complexes can hydrolyze to tocopherylquinone. Of less importance are interactions between
Figure 3  The different resonance structures of the α-tocopherol radical. (Adapted from Ref. 6.)
tocopheroxyl and peroxyl radicals, which form an addition product ortho to the phenoxy oxygen followed by elimination of an alkoxy radical, addition of oxygen, and abstraction of a hydrogen to form two isomers of epoxy-8α-hydroperoxytocopherone.

Subsequent hydrolysis leads to the formation of epoxyquinones (see Fig. 6 for an example of this reaction) [9,10]. Formation of epoxide derivatives of tocopherol represents no net reduction of radicals (because an alkoxy radical forms) and a loss of tocopherol from the system, whereas any tocopherylquinone that is formed can be regenerated back to tocopherol in the presence of reducing agents (e.g., ascorbic acid and glutathione; see Sec. VII: Antioxidant Interactions). An additional reaction that can occur is the interaction of two tocopheroxyl radicals to form tocopherol dimers [11].

**B. Synthetic Phenolics**

Phenolic antioxidants for use in foods include synthetic compounds (Fig. 7). Synthetic phenolic antioxidants exhibit varying polarity, with butylated hydroxytoluene (most nonpolar) > butylated hydroxyanisole > tertiary butylhydroquinone > propyl...
Figure 5 Proposed mechanism for the formulation of α-tocopherylquinone from the interaction of a α-tocopherol radical and a phospholipid peroxyl radical. (Adapted from Ref. 4.)
Figure 6  Formation of an epoxyquinone and an alkoxyl radical from the interaction of an \( \alpha \)-tocopherol radical with a peroxyl radical. (Adapted from Ref. 4.)
Figure 7  Structures of several important phenolic free radical scavengers used in foods.
gallate. The antioxidant mechanism of the synthetic phenolics involves the formation of a resonance-stabilized phenolic radical that neither rapidly catalyzes the oxidation of other molecules nor reacts with oxygen to form antioxidant peroxides that autoxidize [5].

Synthetic phenolic radicals can potentially react with each other by means of mechanisms similar to that of \( \alpha \)-tocopherol. These include reactions of two phenolic radicals to form a hydroquinone and a regenerated phenolic, as well as the formation of phenolic dimers. The phenolic radicals can also react with other peroxyl radicals in termination reactions resulting in formation of phenolic--peroxyl species adducts. In addition, oxidized synthetic phenolic undergo numerous degradation reactions (for review, see Ref. 5). Since many of these degradation products still contain active hydroxyl groups, the products may retain antioxidant activity. Therefore, the net antioxidant activity of synthetic phenolics in food actually represents the activity of the original phenolic plus some of its degradation products. Synthetic phenolics are effective in numerous food systems; however, their use in the food industry has recently declined, reflecting safety concerns and consumer demand for all-natural products.

C. Ubiquinone

Ubiquinone, or coenzyme Q, is a phenolic conjugated to an isoprenoid chain. Ubiquinone is primarily associated with the mitochondrial membrane [12]. Reduced ubiquinone is capable of inactivating peroxyl radicals, but its radical scavenging activity is less than that of \( \alpha \)-tocopherol [13]. The lower free radical scavenging activity of reduced ubiquinone has been attributed to internal hydrogen bonding, which makes hydrogen abstraction more difficult [13]. Despite its lower radical scavenging activity, reduced ubiquinone has been found to inhibit lipid oxidation in liposomes [14] and low density lipoprotein [15]. Presumably, it could be an important endogenous antioxidant in many foods.

D. Plant Phenolics

Plants contain a diverse group of phenolic compounds including simple phenolics, phenolic acids, anthocyanins, hydrocinnamic acid derivatives, and flavonoids. Widely distributed in plant foods such as fruits, spices, tea, coffee, seeds, and grains, these phenolics have been estimated to be consumed in amounts greater than 1 g/day.

All the phenolic classes have the structural requirements of free radical scavengers (see Fig. 7 for several examples). However, the antioxidant activity of these compounds varies greatly, and some even exhibit prooxidant activity. Factors influencing the antioxidant activity of plant phenolics include position and degree of hydroxylation, polarity, solubility, reducing potential, stability of the phenolic to food processing operations, and stability of the phenolic radical. In addition, many phenolics contain acid or ring groups that may participate in metal chelation. These metal chelation properties, in addition to high reducing potentials, can accelerate metal-catalyzed oxidative reactions, leading to the prooxidative activity of plant phenolics under certain conditions [16,17].

Herbs and spices are sources of phenolic antioxidants used in foods. Rosemary extracts are the most commercially important source of an antioxidant ingredient.
containing plant phenolics. Carnosic acid, carnosol, and rosmarinic acid are the major antioxidant phenolics in rosemary extracts (Fig. 7) [18]. Crude rosemary extracts have been found to inhibit lipid oxidation in a wide variety of food products including meats, bulk oils, and lipid emulsions [18–20]. Utilization of phenolic antioxidants from crude herb extracts such as rosemary is often limited by the presence of highly flavorful monoterpenes. Use of more purified forms of herbal phenolics is restricted by both economic and regulatory hurdles.

E. Carotenoids

Carotenoids are a diverse group (>600 compounds) of yellow to red polyenes consisting of 3 to 13 double bonds and in some cases 6 carbon hydroxylated ring structures at one or both ends of the molecule [21]. Carotenoids may be important biological antioxidants and are thought to play a role in controlling oxidatively induced diseases such as cancer and atherosclerosis [22]. The antioxidant properties of carotenoids depend on environmental conditions and the nature of oxidation catalyst. Carotenoids can be effective antioxidants in the presence of singlet oxygen (see Sec. III-A: Control of Prooxidant Metals). However, when peroxyl radicals are the initiating species, the antioxidant efficiency of carotenoids depends on oxygen concentrations.

β-Carotene, the most extensively studied carotenoid antioxidant, reacts with lipid peroxyl radicals, resulting in the formation of a carotene radical. Burton and Ingold [23] found that under conditions of high oxygen tension, the antioxidant activity of β-carotene is diminished. They proposed that increasing oxygen results in increased formation of carotenoid peroxyl radicals, thus favoring autoxidation of β-carotene over inactivation of lipid peroxyl radicals. Under conditions of low oxygen tension, the lifetime of the carotenoid radical is long enough to permit reaction with another peroxyl radical, thus forming a nonradical species and effectively inhibiting oxidation by removing radicals from the system.

Incubation of β-carotene with peroxyl radical generators in organic solvents at high (atmospheric) oxygen tensions leads to addition reactions to form carotenoid–peroxyl adducts (Fig. 8). Addition of a peroxyl radical to the cyclic end group or the polyene chain followed by loss of an alkoxyl radical leads to the formation of 5,6- and 15,15′-epoxides. Elimination of the alkoxyl radical from the 15,15′ positions can also cause cleavage of the polyene chain, resulting in formation of aldehydes. Since the formation of β-carotene epoxides from the addition of peroxyl radicals results in the formation of an alkoxyl radical, the net change in radical number is zero; thus an antioxidant effect is not expected [24].

β-Carotene is capable of donating an electron to peroxyl radicals to produce a β-carotene cation radical and a peroxyl anion. The β-carotene cation radical is resonance stabilized and does not readily react with oxygen to form peroxides. However, the β-carotene cation radical appears to be strong enough to oxidize other lipophilic hydrogen donors, including tocopherols and ubiquinone [24]. Additional research is needed to identify the oxidation products that form from carotenoids under low oxygen partial pressures. Identification of these products may help determine the exact mechanism by which carotenoids act as free radical scavengers when oxygen concentrations are low. Such knowledge would make it easier to predict when carotenoids will exhibit antioxidant activity.
Figure 8  Products formed from the oxidation of β-carotene by a peroxyl radical. (Adapted from Ref. 24.)

F. Water-Soluble Free Radical Inactivators

Free radicals are generated in the water phase of foods by processes such as the Fenton reaction, which produces hydroxyl radicals from hydrogen peroxide [25,26]. Since free radicals are found in the aqueous phase, biological systems contain water-soluble compounds capable of free radical inactivation. Ascorbic acid and glutathione scavenge free radicals, resulting in the formation of low energy ascorbate and glutathione radicals [3]. While ascorbate and glutathione form low energy radicals, other factors influence whether these compounds will act as antioxidants. Both ascorbate and glutathione are strong reducing compounds. Ascorbate, and in some cases glutathione, will catalyze the reduction of transition metals, which in turn can react with
hydrogen and lipid peroxides to form radicals [27, 28]. Ascorbate also causes the release of iron, which is sequestered to proteins such as ferritin [29]. Therefore, ascorbate and glutathione can potentially exhibit prooxidative activity in the presence of free transition metals or iron-binding proteins. In addition, in the presence of oxygen, glutathione radicals are capable of forming high energy peroxides that can potentially catalyze the oxidation of lipids [3].

Thiols besides glutathione can inactivate free radicals. Cysteine is capable of scavenging free radicals. The energy of the resulting thio radical is high, however, suggesting that it may promote oxidation [3]. Thiocysteic acid is another thiol that can inactivate peroxyl radicals [30]. However, the reduced state of thiocysteic acid, dihydrolipoic acid [31] and cysteine [32], can be prooxidative because their reducing potential, and thus their ability to stimulate metal-catalyzed oxidation, is strong.

Several nitrogenous compounds can inactivate free radicals. Uric acid inactivates both hydroxyl and lipid radicals and inhibits lipid oxidation at physiological concentrations [33]. Uric acid is an important antioxidant in blood plasma [33–35]. Since uric acid is produced in postmortem skeletal muscle via ATP metabolism [36], it might possibly serve as an active endogenous antioxidant in muscle foods.

Amino acids, peptides, and proteins can interact with free radicals. Amino acids, including histidine, tyrosine, phenylalanine, tryptophan, cysteine, proline, and lysine, are capable of inactivating free radicals [37–40]. Blood proteins have been estimated to provide 10 to 50% of the peroxyl radical trapping activity of plasma [34,41]. Serum albumin scavenges carbon-based free radicals partially through the involvement of its free sulfhydryl groups [42]. Amino acids, peptides, and proteins have been reported to inhibit lipid oxidation in bulk and emulsified lipid systems as well as in food products [43–47].

While protein, peptides, and amino acids often possess the structural characteristics needed to both scavenge radicals and inhibit lipid oxidation, the concentrations required for activity are often higher than other free radical scavengers. This suggests that free radicals inactivated by proteinaceous compounds do not act as chain-breaking antioxidants but instead are simply competing with the lipid for high energy radicals. Interactions of free radicals with amino acids and proteins leads to the formation of peroxides [37,48]. However, very little is known about how the formation of amino acid peroxides or other amino acid oxidation products influences the antioxidant activity of amino acids, peptides, and proteins.

III. CONTROL OF LIPID OXIDATION CATALYSTS

Lipid oxidation rates in foods often depend on catalyst concentrations and activity. Control of lipid oxidation catalysts can therefore be a very important factor in controlling oxidative rancidity. Both endogenous and added antioxidants help control the activity of transition metals, singlet oxygen, and enzymes.

A. Control of Prooxidant Metals

Transition metals accelerate lipid oxidation reactions by hydrogen abstraction and peroxide decomposition, resulting in the formation of free radicals [26]. The activity of prooxidative metals is influenced by chelators or sequestering agents. Transition metals such as iron exhibit low solubility at pH values near neutrality [25]. Therefore,
in food systems, transition metals often exist chelated to other compounds. Many compounds will form complexes with metals, resulting in changes in catalytic activity. Some metal chelators increase oxidative reactions by increasing metal solubility and/or altering the redox potential [49]. Chelators also increase the prooxidant activity of transition metal activity by making them more nonpolar, thereby increasing their solubility in lipids [50]. Chelators that exhibit antioxidative properties inhibit metal-catalyzed reactions by one or more of the following properties: prevention of metal redox cycling, occupation of all metal coordination sites, formation of insoluble metal complexes, and steric hindrance of interactions between metals and lipids or oxidation intermediates (e.g., peroxides) [51]. The prooxidative/antioxidative properties can depend on both metal and chelator concentrations. For instance, EDTA is prooxidative when ratios of EDTA to iron are 1 or less and antioxidative when EDTA/iron ≥ 1 [49].

The most common metal chelators used in foods contain multiple carboxylic acids (e.g., EDTA and citric acid) or phosphate (e.g., polyphosphates and phytate) groups. Chelators are typically water soluble but some will exhibit solubility in lipids (e.g., citric acid), thus allowing the chelator to inactivate metals in the lipid phase [52]. Chelator activity depends on pH, since the chelator must be ionized to be active. Therefore, as pH approaches the pK of the ionizable groups, chelator activity decreases. Chelator activity is also decreased by the presence of other chelatable ions (e.g., calcium), which will compete with the prooxidative metals for binding sites.

Although most food-grade chelators are unaffected by food processing operations and subsequent storage, polyphosphates are an exception. Polyphosphates are stronger chelators and antioxidants than mono- and diphosphates [53]. However, some foods contain phosphatases, which hydrolyze polyphosphates, thus decreasing their antioxidative effectiveness. This can be observed in muscle foods, where polyphosphates are relatively ineffective in raw meats that contain high levels of phosphatase activity [54] but are highly effective in cooked meats, where the phosphatases have been inactivated [55]. Nutritional implications should also be considered when chelators are used as food antioxidants, since chelators influence mineral bioavailability. For instance, EDTA enhances iron bioavailability while phytate decreases iron, calcium, and zinc absorption [56].

Prooxidant metal activity is also controlled in biological systems by proteins. Proteins with strong binding sites include transferrin, ovotransferrin (conalbumin), lactoferrin, and ferritin. Transferrin, ovotransferrin, and lactoferrin are structurally similar proteins consisting of a single polypeptide chain with a molecular weight ranging from 76,000 to 80,000. Transferrin and lactoferrin bind two ferric ions apiece, while ovotransferrin has been reported to bind three [29,57,58]. Ferritin is a multisubunit protein (molecular weight 450,000) with the capability of storing up to 4500 ferric ions [59]. Transferrin, ovotransferrin, lactoferrin, and ferritin inhibit iron-catalyzed lipid oxidation by binding iron in its inactive ferric state and possibly by sterically hindering metal–peroxide interactions [29,60]. Reducing agents (ascorbate, cysteine, superoxide anion) and low pH can cause the release of iron from the proteins, resulting in an acceleration of lipid oxidation reactions [29,61]. The activity of copper can also be controlled by binding to proteins. Serum albumin binds one cupric ion [62] and ceruloplasmin binds up to six cupric ions [63].

Amino acids and peptides can chelate metals in a manner that decreases their reactivity. Both the chelating and the antioxidant activities of the skeletal muscle.
dipeptide carnosine depend on metal ion type [64–66]. Carnosine more effectively inhibits the oxidation of phosphatidylcholine liposomes catalyzed by copper than by iron. Decker et al. [66] found that the carnosine can chelate and inhibit the prooxidant activity of copper but more effectively than its constituent amino acid histidine.

Ceruloplasmin is a copper-containing enzyme that catalyzes the oxidation of ferrous ions:

\[ 4\text{Fe}^{2+} + 4\text{H}^+ + \text{O}_2 \rightarrow 4\text{Fe}^{3+} + 2\text{H}_2\text{O} \]

This ferroxidase activity inhibits lipid oxidation by maintaining iron in its oxidized, inactive state [63]. Since ceruloplasmin is primarily a constituent of blood, one would not expect to find it in most foods, other than muscle foods. Addition of ceruloplasmin to muscle foods in a pure form or as part of blood plasma has been found to effectively inhibit lipid oxidation [67,68].

B. Control of Singlet Oxygen

Singlet oxygen is an excited state of oxygen in which two electrons in the outer orbitals have opposite spin directions. Initiation of lipid oxidation by singlet oxygen is due to its electrophilic nature, which leads to the formation of lipid peroxides from unsaturated fatty acids [69].

Singlet oxygen can be inactivated by both chemical and physical quenching. Chemical quenching of singlet oxygen by \( \beta \)-carotene will lead to the formation of carotenoid breakdown products containing aldehyde and ketone groups as well as \( \beta \)-carotene-5,8-endoperoxide. \( \beta \)-Carotene-5,8-endoperoxide, which occurs mainly upon the oxidation of \( \beta \)-carotene by singlet oxygen, therefore may provide a unique marker that could be used to monitor singlet oxygen/carotenoid interactions in foods and biological systems [70]. Tocopherols can chemically quench singlet oxygen in reactions that lead to the formation of tocopherol peroxides and epoxides [69]. Other compounds, including amino acids, peptides, proteins, phenolics, urate, and ascorbate, can chemically quench singlet oxygen, but little is known about the resulting oxidation products [69,71,72].

While carotenoids are capable of chemically inactivating singlet oxygen, these reactions cause carotenoid autoxidation, leading to loss of antioxidant activity. Therefore, the major mechanism of singlet oxygen inactivation by carotenoids is physical quenching. The most common energy states of singlet oxygen are 22.4 and 37.5 kcal above the ground state [69]. Carotenoids physically quench singlet oxygen by a transfer of energy to the carotenoid to produce an excited state of the carotenoid and ground state, triplet oxygen. Energy is dissipated from the excited carotenoid by vibrational and rotational interactions with the surrounding solvent to return the carotenoid to the ground state [73]. Nine or more conjugated double bonds are necessary for physical quenching [74]. The presence of six carbon-oxygenated ring structures at the end of the polyenes increases the ability of carotenoids to physically quench singlet oxygen [74]. While it is generally believed that the physical quenching of singlet oxygen by carotenoids does not cause destruction of the carotenoid, these reactions may result in trans or cis isomer conversions [73].

Tocopherols and amines can physically quench singlet oxygen by a charge transfer mechanism. In this reaction, the electron donor (tocopherol or amine) forms a charge transfer complex with the electron-deficient singlet oxygen molecule [70].
An intersystem energy transfer occurs in the complex, resulting in a dissipation of energy and the eventual release of triplet oxygen.

C. Control of Lipoxygenases

Lipoxygenases are active lipid oxidation catalysts found in plants and some animal tissues. Lipoxygenase activity can be controlled by heat inactivation and plant breeding programs that decrease the concentrations of these enzymes. Phenolics are capable of indirectly inhibiting lipoxygenase activity by acting as free radical inhibitors, but also by reducing the iron in the active site of the enzyme to the catalytically inactive ferrous state [16].

IV. INACTIVATION OF OXIDATION INTERMEDIATES

Several compounds that can exist in foods indirectly influence lipid oxidation rates. While these substances do not always directly interact with lipids, they may interact with metals or oxygen to form reactive species. Examples of such compounds include superoxide anion, peroxides, and photosensitizers.

A. Superoxide Anion

Superoxide anion is produced by the addition of an electron to molecular oxygen. Superoxide participates in oxidative reactions by maintaining transition metals in their reduced, active states, by promoting the release of metals bound to proteins such as ferritin, and through the pH-dependent formation of its conjugated acid, the perhydroxyl radical, which can directly catalyze lipid oxidation [26]. Because superoxide anion participates in oxidative reactions, biological systems contain superoxide dismutase (SOD).

Two forms of SOD are found in eukaryotic cells, one in the cytosol and the other in the mitochondria [75]. Cytosolic SOD contains copper and zinc in the active site, while mitochondrial SOD contains manganese. Both forms of SOD catalyze the conversion of superoxide anion to hydrogen peroxide by the following reaction:

\[ 2O_2^- + 2H^+ \rightarrow O_2 + H_2O_2 \]

Other compounds can also possess superoxide dismutase-like activity. The most notable of these are complexes of amino acids and peptides with transition metals. Cupric ions complexed to lysine, tyrosine, and histidine are capable of catalyzing the dismutation of superoxide [76]. Histidine-containing peptides complexed to nickel [77], copper [78], and zinc [78,79] also contain superoxide dismutase-like activity. It should be noted that the SOD-like activity of metal—amino acid or peptide complexes generally is orders of magnitude lower than those of proteinaceous SOD.

B. Peroxides

Peroxides are important intermediates of oxidative reactions because they decompose via transition metals, irradiation, and elevated temperatures to form free radicals. Hydrogen peroxide exists in foods as a result of direct addition (e.g., aseptic processing operations) and formation in biological tissues by mechanisms including the dismutation of superoxide by SOD and the activity of peroxisomes [84]. Hydrogen
peroxide is rapidly decomposed by the reduced state of transition metals (e.g., Fe and Cu) to the hydroxyl radical. The hydroxyl radical is an extremely reactive free radical that can oxidize most biological molecules at diffusion-limited reaction rates. Therefore, removal of hydrogen peroxide from biological materials is critical to the prevention of oxidative damage.

Catalase (CAT) is a heme-containing enzyme that catalyzes the following reaction [81]:

$$2\text{H}_2\text{O}_2 \rightarrow 2\text{H}_2\text{O} + \text{O}_2$$

Hydrogen peroxide in higher plants and algae may be scavenged by ascorbate peroxidase. Ascorbate peroxidase inactivates hydrogen peroxide in the cytosol and chloroplasts by the following mechanism [82]:

$$2\text{ascorbate} + \text{H}_2\text{O}_2 \rightarrow 2\text{monodehydroascorbate} + 2\text{H}_2\text{O}$$

Two ascorbate peroxidase isozymes which differ in molecular weight (57,000 vs. 34,000), substrate specificity, pH optimum, and stability have been described in tea leaves [83].

In addition to catalase, many biological tissues contain glutathione peroxidase (GSH-Px) to help control peroxides. GSH-Px differs from CAT in that it is capable of reacting with both lipid and hydrogen peroxides. GSH-Px is a selenium-containing enzyme that uses reduced glutathione (GSH) to catalyze hydrogen or lipid (LOOH) peroxide reduction [84]:

$$\text{H}_2\text{O}_2 + 2\text{GSH} \rightarrow 2\text{H}_2\text{O} + \text{GSSG}$$

or

$$\text{LOOH} + 2\text{GSH} \rightarrow \text{LOH} + \text{H}_2\text{O} + \text{GSSG}$$

where GSSG is oxidized glutathione and LOH is a fatty acid alcohol. Two types of GSH-Px exist in biological tissues, and one shows high specificity for phospholipid hydroperoxides [84,85].

Thiodipropionic acid and dilauryl thiodipropionate are approved food additives capable of decomposing peroxides and peracids. At the concentrations allowed in foods (≤200 ppm); however, they are relatively ineffective antioxidants and are therefore rarely used [52]. Methionine, which has been found to be antioxidative in some lipid systems, is thought to decompose peroxides by mechanisms similar to those of thiodipropionic acid and dilauryl thiodipropionate [52].

C. Photoactivated Sensitizers

In foods, light is capable of activating sensitizers such as chlorophyll, riboflavin, and heme-containing proteins to an excited state. These photoactivated sensitizers can promote oxidation by directly interacting with an oxidizable substrate to produce free radicals, by transferring energy to triplet oxygen to form singlet oxygen, or by transfer of an electron to triplet oxygen to form superoxide anion [69]. Carotenoids inactivate photoactivated sensitizers by physically absorbing their energy to form the excited state of the carotenoid, which then returns to the ground state by transfer of energy into the surrounding solvent [22,73].
V. ALTERATIONS IN LIPID OXIDATION BREAKDOWN PRODUCTS

Oxidation of fatty acids eventually leads to formation of breakdown products via $\beta$-scission reactions. These reactions lead to a multitude of different oxidation products, known as secondary lipid oxidation products, which affect both the sensory characteristics and the functional properties of foods. Rancid odors arise from the production of secondary products such as aldehydes, ketones, and alcohols [86]. Secondary lipid oxidation products, and in particular aldehydes, also impact food quality and nutritional composition through interaction with the amino groups of proteins and vitamins. Secondary products arising from lipid oxidation have been found to alter the function of proteins, enzymes, biological membranes, lipoproteins, and DNA [80,87–89].

Since aldehydes and other secondary products arising from lipid oxidation are potentially damaging, biological systems seem to have developed mechanisms to control their activity. Sulfur- and amine-containing compounds have the ability to interact with aldehydes. This may help explain why many proteins, peptides, amino acids, phospholipids, and nucleotides display antioxidant activity when secondary products are used to measure lipids oxidation. Carnosine and anserine, which can make up over 1% of the wet weight of muscle tissue, are capable of forming complexes with aldehydes produced from oxidizing lipids [90]. Carnosine is more effective at forming adducts with aldehydes than its constituent amino acids, histidine and $\beta$-alanine [91,92]. Glutathione is also very effective at binding aldehydes, but at the concentrations found in muscle foods, carnosine seems more likely to be the major aldehyde-binding component [92].

VI. SURFACE-ACTIVE ANTIOXIDANTS AND PHYSICAL EFFECTS

Lipids in food systems often have interfacial surfaces at which oxidative reactions are prevalent. Examples include oil-in-water emulsions, water-in-oil emulsions, the air–lipid interface of bulk oils and solid fats, and the water–lipid interface of biological membranes. Oxidation is prevalent at these interfaces as a result of increased contact with oxygen, the presence of aqueous phase free radicals, the presence of reactive oxygen generating systems and prooxidative metals, and possibly the migration of the more polar lipid peroxides out of the hydrophobic lipid core toward the more polar interface.

The effectiveness of phenolic antioxidants is often dependent on their polarity. Porter [93] used the term “antioxidant paradox” to describe how polar antioxidants are most effective in bulk lipids while nonpolar antioxidants are most effective in dispersed lipids. In bulk tocopherol-stripped corn oil, Trolox (a water-soluble analog of $\alpha$-tocopherol) more effectively inhibited lipid peroxide formation than $\alpha$-tocopherol. However, when tocopherol-stripped corn oil was emulsified with Tween 20, $\alpha$-tocopherol inhibited peroxide formation more effectively than Trolox. The observed increase in activity of $\alpha$-tocopherol compared to Trolox in emulsified oil was attributed to its retention in the oil and possibly to its ability (due to its surface activity) to concentrate at the oil–water interface. The lower activity of Trolox in emulsions was due to its partitioning into the water phase, where it was not able to inhibit autoxidation of the corn oil [94]. Similar effects have been observed for the phenolic antioxidants in rosemary extracts, with the more polar compounds (carnosic and
Rosmarinic acids) being most effective in bulk oils and the less polar compounds (carnosol) being more effective in emulsified lipids [18]. Similarly, the antioxidant activity of carnosic acid was improved in emulsified corn oil when it was made nonpolar by methylation [95].

The charge of dispersed lipids also influences oxidation rates, especially in the presence of transition metals. Since iron and other transition metals are common contaminants in most water systems, their ability to catalyze oxidation at the oil–water interface of dispersed lipids could be important. When the surface charge of dispersed lipids as either micelles [96,97] or phospholipid vesicles [98] is negative, iron-catalyzed lipid oxidation rates are much higher than they are at positively charged interfaces. This effect presumably exists because iron can bind to the interface of the dispersed lipid. The inhibitory effect of positively charged lipid micelles can be partially overcome by nitrilotriacetic acid, which forms negatively charged iron chelates, and by addition of lipid-soluble peroxides [96,97]. Positively charged emulsifiers are uncommon in foods. However, proteins at pHs below their pIs could produce positively charged lipid emulsion droplets. Unfortunately, little is known about the ability of proteins to influence oxidation by modifying the charges that govern metal attraction/repulsion. The charge of dispersed lipid droplets can also be important for the activity of antioxidants. Negatively charged ascorbic acid is a more effective antioxidant in the presence of a positively charged lipid dispersion [99], while spermine (positively charged) is more effective when the lipid interface is positive [100].

Lipid oxidation can be inhibited by encapsulation. Potential mechanisms of inhibition include physical inhibition of oxygen diffusion into the lipid, chemical (e.g., free radical scavenging) and physical (e.g., chelation) antioxidant properties of the encapsulating agents, and possibly interaction of lipid oxidation products with the encapsulating material. Both protein and carbohydrate encapsulating agents have been found to retard oxidation rates. The effectiveness of these encapsulating agents depends on factors such as concentration of the encapsulating agent [101], method of encapsulation (which affects the porosity of the encapsulating layer) [102], and the environmental relative humidity under which the encapsulated lipid is stored [103].

Some evidence indicates that encapsulation does inhibit oxygen diffusion into the lipid, but the same research also indicates that oxygen diffusion is not the only mechanism by which encapsulation inhibits oxidation [104]. More research is needed to determine the antioxidant mechanisms of encapsulation, since this technique could be an effective way to increase the stability of oxidatively labile lipids, thereby increasing their incorporation into foods.

Another factor that may influence oxidation rates is the physical state of the lipid. Lipids in foods often exist as a combination of both liquid and crystalline states, a condition that depends on both fatty acid composition and temperature. The influence of liquid fat concentration on oxidation rates was investigated in liposomes, where arachidonic acid oxidation rates were found to be greater at temperatures below the solid–liquid phase transition temperature of the host lipid [105]. The increase in oxidation rates was attributed to phase separation of the most unsaturated fatty acids, which increases the concentrations of oxidizable substrate into localized domains [106]. Little is known about how transition temperatures influence oxidation rates in food lipids.
VII. ANTIOXIDANT INTERACTIONS

Biological food systems usually contain multicomponent antioxidant systems. The numerous existing antioxidants have different potential functions, including inhibition of prooxidants of different types (e.g., metals, reactive oxygen species, enzymes); inactivation of free radicals and prooxidants in aqueous, interfacial, and lipid phases; and inactivation of compounds at different stages of oxidation [e.g., initiating species (·OH), propagating species (peroxides), lipid oxidation decomposition products (aldehydes)]. In addition, multicomponent antioxidant systems are beneficial because direct interactions occur between antioxidants.

Combinations of chelators and FRS often result in synergistic inhibition of lipid oxidation [6]. Synergistic interaction most likely occurs by a “sparking” effect provided by the chelator. Since the chelator will decrease oxidation rates by inhibiting metal-catalyzed oxidation, fewer free radicals will be generated in the system. This means that the eventual inactivation of the FRS through reactions such as termination or autoxidation will be slower, thus making its concentration greater at any given time. The combination of chelator and FRS thus decreases free radical generation and increases radical scavenging potential.

Synergistic antioxidant activity can also be observed by the combination of two or more different FRS. This occurs when one FRS reacts more rapidly with free radicals than the other as a result of differences in bond disassociation energies and/or steric hindrance of FRS/ROO· interactions [6]. These differences will result in one antioxidant being consumed faster than the other. However, it may be possible for this FRS to be regenerated by transfer of its radical to a different scavenger. In the system consisting of α-tocopherol and ascorbic acid [106], for example, α-tocopherol is the primary FRS because it is present in the lipid phase. Ascorbic acid then regenerates the tocopheroxyl radical or possibly tocopherylquinone back to α-tocopherol plus the semihydroascorbic radical [4], which dismutates to dehydroascorbate [3]. In turn, dehydroascorbate may be regenerated by enzymes that utilize NADH or NADPH as reducing equivalents [107].

Since multicomponent antioxidant systems can inhibit oxidation at many different phases of oxidation, the resulting antioxidant activity can be synergistic. This suggests that the most effective antioxidant systems for foods would contain antioxidants with different mechanisms of action and/or physical properties. Determining which antioxidants would be most effective depends on factors such as types of oxidation catalyst, physical state of lipid (bulk vs. emulsified), and factors that influence the activity of the antioxidants themselves (e.g., pH, temperature, the ability to interact with other compounds in the foods).

REFERENCES


I. FATS AND OILS IN HUMAN HEALTH

Most discussions of fats and health focus on the deleterious effects of these essential nutrients. What we are really discussing in that case is the possibly harmful effects of an excess of fats and oils. Fats (lipids) supply energy, support structural aspects of the body, and provide substances that regulate physiological processes.

Adipose tissue, which is the repository of most of our body fats, serves as an energy reservoir (fat supplies nine calories per gram compared to four calories per gram for protein or carbohydrate), as a heat conserver and as a shock absorber. Lipids contain essential fatty acids, such as linoleic and linolenic acids. These are metabolized eventually to provide eicosanoids, substances that possess hormone-like activity and thus may regulate many body functions. Fat is also the transport vehicle for vitamins A, D, E, and K.

Cholesterol, which has absorbed the brunt of the antifat attack, is a compound that is essential for life. It is not essential in the sense of essential fatty acids since the body can synthesize it, but it is a crucially important component of our biological economy. Cholesterol comprises about 0.2% of normal body weight. Most of it (about 33%) is in the brain and nervous system where its function has not been probed beyond suggesting that its major function is as an insulator. Almost another one-third of the body’s cholesterol is in muscle where it is a structural component. Every cell membrane contains cholesterol and phospholipid, another fatty substance. The esterified cholesterol found in muscle may represent a storage compartment. The percentage of cholesterol ester in muscle increases with age. Cholesterol is the parent substance for vitamin D₃, bile acids, adrenocortical hormones, and sex hormones. Thus, it is one of the more important biological substances. Fat also contributes to the palatability and flavor of food and hence contributes to the enjoyment of eating.
The two major causes of death in the developed world are heart disease and cancer. Both have been described as lifestyle diseases, and effects of diet fit under that rubric. Since diet is one of the easiest lifestyle factors to investigate and possibly change, its role has been pursued with vigor. However, dietary data are not as simple to obtain as one might expect. Population-based data, derived from availability statistics, do not account for individual variations. Recall may be flawed by habitual underreporting of intake [1,2]. These methods provide useful data but their shortcomings should be kept in mind.

II. LIPIDS AND CARDIOVASCULAR DISEASE

The major difficulty in assessment of the roles of diet and other factors is the absence of a clear, unequivocal, antemortem diagnosis. Failing that, the data have been analyzed to provide “risk” factors (called “odds” in Las Vegas) for indication of susceptibility. Among the major risk factors for coronary heart disease (CHD), also termed cardiovascular disease (CVD), are cigarette smoking, elevated cholesterol level, elevated blood pressure, obesity, and maleness. The first of these is correctable and the last is an unalterable fact of life, but the others may, to some extent, be amenable to nutritional intervention. However, it is oversimplification to regard atherosclerosis as a consequence of diet or aging or both. The molecular mechanisms underlying the atherosclerotic process are being elucidated, and increasingly aspects of the disease exhibit a genetic component. We are discovering new molecular and immunological factors related to this disease. Fatal outcome is associated with plaque rupture and thrombosis, and the notion of the disease being due to simple accretion of cholesterol in the arteries is no longer tenable. Lusis [3] has published an elegant description of factors involved in atherogenesis.

Although an experimental relationship between dietary cholesterol and atherosclerosis had been adduced in 1913 when Anitschkow was able to produce aortic lesions in rabbits [4], interest in fat in the diet and its relation to this disease began to blossom in the 1950s. In 1950 Gofman and his colleagues [5] developed a method for the ultracentrifugal separation of plasma lipoproteins, showed how these fractionated particles could be related to heart disease, and implicated diet as a factor. In 1952 Keys [6] described what was the beginning of his Seven Countries Study and indicated atherosclerosis as a new public health problem.

The lipoproteins are lipid–protein agglomerates rather than real chemical compounds. They are described by their hydrated densities (a physical property) but may differ in size and composition. Thus, although the chemical analyses of low density lipoprotein (LDL), high density lipoprotein (HDL), etc., are often published, they represent average values and are not as precise indicators of identity as melting point or spectrum. As research continues we continue to discover subfractions of lipoproteins that affect CHD risk. Lipoprotein (a) [Lp(a)], which was described first by Berg [7], is an LDL particle in which apoprotein B is linked to an apoprotein unit [apo-protein (a)] via a disulfide bridge. Elevated Lp(a) levels have been associated with a high risk of CHD [8–10]. Lp(a) interferes with fibrinolysis [11], and its levels in the blood appear not to be affected by diet [12] or drugs [13].

Krauss and his co-workers [14,15] have identified subpopulations of LDL particles that differ in size and composition (Table 1). The smaller, denser particles may be associated more strongly with the risk of coronary disease [16–18], and their
Table 1  Classification of LDL Particles

<table>
<thead>
<tr>
<th>Class</th>
<th>Subfraction density (g/mL)</th>
<th>Particle diameter (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDL I</td>
<td>1.025–1.035</td>
<td>26–27</td>
</tr>
<tr>
<td>LDL II</td>
<td>1.032–1.038</td>
<td>25.5–26</td>
</tr>
<tr>
<td>LDL IIIA</td>
<td>1.038–1.050</td>
<td>24.7–25.6</td>
</tr>
<tr>
<td>LDL IIIB</td>
<td>24.2–24.6</td>
<td></td>
</tr>
<tr>
<td>LDL IV A</td>
<td>1.048–1.065</td>
<td>23.3–24.2</td>
</tr>
<tr>
<td>LDL IVB</td>
<td>21.8–23.2</td>
<td></td>
</tr>
</tbody>
</table>

Note  LDL receptor activity and antioxidant content highest in LDL I and II. Triglyceride content increases with decreasing size.

Source: From Ref. 15.

levels may be determined genetically [19]. Animal studies had shown earlier that large lipoprotein molecules do not enter the arterial wall [20,21].

Cholesterol has assumed a central role in experimental and human atherosclerosis, and the public is being exhorted to know its “cholesterol number.” However, cholesterol levels tend to vary diurnally and with season [22–24], and single measurements may not be an accurate indicator of risk. This is especially true if the single determined value is near one of the accepted cut points [25]. Low levels of cholesterol may lead to increased risk of noncardiovascular death [26–28]. Low cholesterol may become a problem at levels below 160 mg/dL [28] or 180 mg/dL [26].

Since ingested cholesterol has been shown to be atherogenic in some animal species, since elevated levels of cholesterol are a risk factor, and since it is relatively easy to measure, cholesterol has borne the brunt of the attack on CHD. The effect of dietary cholesterol on levels of blood cholesterol appears to be small. In 1950 Gertler et al. [29] isolated from a large study of coronary disease 4 groups of 10 men each. They were the men with lowest or highest serum cholesterol and those who ingested the most or the least dietary cholesterol. They were compared with similar groups selected from the control subjects. In every group the men with coronary disease exhibited significantly higher cholesterol levels than the controls but in no case was any relation to cholesterol intake seen. Early in the Framingham Study it was found that plasma cholesterol levels were unrelated to diet [30], a finding also reported from the Tecumseh Study [31]. Several groups have reported that addition of eggs to the diet of free-living subjects did not affect their cholesterol level [32–34]. Gordon et al. [35] analyzed and compared the diets of men who did or did not have coronary disease in three large prospective coronary disease studies —Framingham, Puerto Rico, and Honolulu. Men who had coronary disease ingested fewer total calories, less carbohydrate, and less alcohol. Intakes of cholesterol and the P/S ratio of their dietary fat were similar for men who did or did not have coronary disease. McNamara [36] reviewed data from 68 clinical studies relating to effect of dietary cholesterol on plasma cholesterol. He concluded that there was a mean rise of 2.3 ± 0.2 mg/dL of plasma cholesterol for every 100 mg of ingested cholesterol. Hopkins [37] described the complexity of the association between cholesterol intake and plasma cholesterol. He found that the magnitude of the change...
in plasma cholesterol as a function of dietary cholesterol is influenced by baseline cholesterol intake. A recent epidemiological overview [38] concluded that after one considered dietary confounders there was no association between egg consumption at levels up to 1 egg per day and the risk of coronary heart disease in nondiabetic men and women.

In contrast to dietary cholesterol there is little question that the saturation of dietary fat exerts a profound influence on blood cholesterol levels. Ahrens et al. [39] fed a number of subjects a liquid formula diet containing 45% of energy as fat. In general, plasma cholesterol rose as the fat saturation rose. Keys et al. [40] and Hegsted et al. [41] studied the effects of changes in dietary fat on change in blood cholesterol levels in humans and offered formulas to predict changes in cholesterolemia based on changes in dietary fat. Both groups found fat saturation to have the greatest effect. Stearic acid did not appear to fit the formula, and direct experiments in human subjects have shown this to be true [42,43]. Hayes and Khosla [44] have hypothesized that the two most important fatty acids related to cholesterol levels are myristic acid (which raises cholesterol levels at every concentration) and linoleic acid (which exerts an increasing hypocholesterolemic effect until it reaches a dietary level of 6–7% of energy). Hayes [45] has reviewed these data recently. McNamara et al. [46] fed normal subjects diets high or low in cholesterol and containing saturated or unsaturated fat. The major factor determining cholesterolemia was the saturation of the fat, the influence of which was about four times greater than that of dietary cholesterol (Table 2). McNamara [47] has reviewed exhaustively the connection between dietary cholesterol and atherosclerosis. He cites large epidemiological studies that indicate little connection between cholesterol intake and risk of coronary heart disease [48,49], emphasizes the role of saturated fat, and suggests that diets very high in cholesterol reflect an unbalance between intake of fats and of grains, vegetables, and fruits.

In addition to fatty acid saturation the position of a specific fatty acid in the triglyceride molecule is important [50]. In an effort to test cholesterolemic effects of specific fatty acids, McGandy et al. [51] fed human subjects diets that contained fats into which high levels of specific saturated fatty acids (lauric, myristic, palmitic, or

### Table 2  Plasma Cholesterol Levels in Subjects Fed High or Low Levels of Cholesterol with Saturated or Unsaturated Fat

<table>
<thead>
<tr>
<th>Fat in diet</th>
<th>Cholesterol</th>
<th>Dietary (mg)</th>
<th>Plasma (mg/dL)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fat (P/S)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low cholesterol</td>
<td>0.31 ± 0.18</td>
<td>288 ± 64</td>
<td>243 ± 50</td>
</tr>
<tr>
<td>Saturated</td>
<td>1.90 ± 0.90</td>
<td>192 ± 60</td>
<td>218 ± 46</td>
</tr>
<tr>
<td>Unsaturated</td>
<td>0.27 ± 0.15</td>
<td>863 ± 161</td>
<td>248 ± 51</td>
</tr>
<tr>
<td>High cholesterol</td>
<td>1.45 ± 0.50</td>
<td>820 ± 102</td>
<td>224 ± 46</td>
</tr>
</tbody>
</table>

*Source: From Ref. 46.*
Table 3 Influence of Native and Randomized Tallow and Lard on Atherosclerosis in Rabbits

<table>
<thead>
<tr>
<th>Group</th>
<th>Tallow</th>
<th>Randomized tallow</th>
<th>Lard</th>
<th>Randomized lard</th>
</tr>
</thead>
<tbody>
<tr>
<td>No.</td>
<td>7/8</td>
<td>7/8</td>
<td>8/8</td>
<td>8/8</td>
</tr>
<tr>
<td>% 16:0 at SN2</td>
<td>3.7(^a)</td>
<td>8.5(^b)</td>
<td>20.8(^b)</td>
<td>7.6(^b)</td>
</tr>
</tbody>
</table>

Plasma
Cholesterol (mg/dL)

| Arch   | 1.29 \(\pm\) 0.24 | 1.50 \(\pm\) 0.53 | 2.69 \(\pm\) 0.28\(^c\) | 1.50 \(\pm\) 0.28\(^c\) |
| Thoracic | 0.79 \(\pm\) 0.26 | 0.79 \(\pm\) 0.28 | 1.75 \(\pm\) 0.28\(^d\) | 0.69 \(\pm\) 0.19\(^d\) |

\(^a\)Of 24.8% total.
\(^b\)Of 21.4% total.
\(^c\)Graded on 0–4 scale.
\(^d\)Lard vs. randomized lard.

Note: Rabbits were fed semipurified diet containing 0.4% cholesterol for 60 days. \((p < 0.05)\).

Source: From Ref. 53.
cholesterolemic effects of trans fats seen to be related to the ratio of dietary trans fat to linoleic acid, the cholesterol levels rising as the ratio falls. Several studies have shown that the tissue levels of *trans*-fatty acids are no higher in subjects with coronary disease than in controls [68,69]. Houtsömüller [70] suggested in 1978 that trans fat be regarded as a quasi-saturated fat.

In 1985 the Life Sciences Research Office of the Federation of American Societies of Experimental Biology published a report which concluded that there was little reason for health concerns at the reported intake level of trans fat (8 g/person/day) [71]. Two years later the British Nutrition Foundation published a report with the same conclusion [72]. A decade later reports sponsored by the International Life Sciences Institute [73] and the British Nutrition Foundation [74] have found no reasons to alter their previous conclusions. All the reports contained the safe suggestion that more research was needed. Two reviews of trans fat effects have appeared recently [75,76]. The findings that trans fats may elevate plasma LDL cholesterol levels, thus increasing risk, suggest that it might be prudent to replace them when possible. It should also be noted that between 1960 and 1985, levels of trans fats in the American diet were fairly constant (7.63 ± 0.08 g/person/day) [77], whereas total age-adjusted mortality and deaths from heart disease and strokes fell by 28%, 37%, and 59%, respectively [78].

Several new players have appeared on the heart disease stage and they may ultimately affect our views of fat and cholesterol as major players in the CHD arena. Over 30 years McCully [79] suggested that homocysteinemia could be a major risk for coronary disease, and this is being recognized today [80–82]. Studies comparing lipid levels in European countries show little relation of these levels to CHD mortality. Ischemic heart disease mortality is four times higher in Belfast, Ireland than in Toulouse, France despite general similarities in their diets (the French ingest significantly more cholesterol and alcohol) and in their risk factor profiles [83]. Similarly, mortality from heart disease in 50- to 54-year-old men is four times higher in Vilnius, Lithuania than it is in Linköping, Sweden despite the fact that differences in traditional risk factors are small [84]. In the years 1985–1987 age-specific mortality from ischemic heart disease for men aged 45–54 years was 237/100,000 in Belfast and 56/100,000 in Toulouse. In men aged 55–64 the rates per 100,000 were 761 in Belfast and 175 in Toulouse. In the younger age group total cholesterol levels were significantly lower in the French (230 ± 41 mg/dL vs. 240 ± 41 mg/dL) and HDL cholesterol levels were higher (54 ± 15 mg/dL vs. 47 ± 12 mg/dL). There were no differences in the older men. Energy intake was virtually the same in the two groups. The French ingested significantly more protein and cholesterol and significantly less carbohydrate while fat intake was similar. In 1977 the CHD mortality per 100,000 men aged 50–54 was 300 in Lithuania and 220 in Sweden. In 1994 mortality was 445 in Lithuania (102% increase) and 110 in Sweden (a 50% decrease). Data from the Russian Lipid Research Clinics show an appreciable number of deaths in men with low levels of LDL and high levels of HDL [85].

The isolation of *Chlamydia pneumoniae* from atherosclerotic, but not normal, arteries [86,87] and the finding of cytomegalovirus in diseased arteries [88] may shed light on mechanisms underlying the onset of the disease.

The role of conjugated linoleic acid (CLA) in atherogenesis is under study. The CLA present in the diet (in dairy products and meat of ruminant animals) is primarily octadeca-c9,t11-dienoic acid, but the commercial product used in most
studies contains equal amounts (40–45%) of the c9,t11, and t10,c12 modifications. CLA has been shown to inhibit atherogenesis in cholesterol-fed rabbits [89] and hamsters [90]. Of greater interest is the observation that feeding 1% CLA to rabbits with preestablished atherosclerosis leads to significant regression [91] of lesions.

III. LIPIDS AND CANCER

In 1930 Watson and Mellanby [92] showed that the incidence of coal tar–induced skin tumors in mice rose from 34% to 57% when 12.5–25.0% butter was added to the basal diet that normally contained 3% fat. Baumann et al. found that high-fat diets increased the yield of ultraviolet radiation–induced or chemically induced skin tumors in mice [93]. They also found saturated fats to be less cocarcinogenic than unsaturated fats [94]. Carroll and Khor [95] made a similar observation. The reason that unsaturated fat enhanced carcinogenesis was found by Ip et al. [96] to be due to the tumor’s requirement for linoleic acid as a growth factor. This finding may explain why rats fed fish oils [97] or fats high in trans unsaturated [98,99] fatty acids also show a reduced incidence of tumors.

Armstrong and Doll [100] published a thorough review correlating cancer incidence in over 30 countries with diet, gross national product (GNP), physician density, population density, and use of solid or liquid fuel. They found positive associations between breast and colorectal cancers and total fat consumption. They also found a strong association between these cancers and GNP. In their conclusion they state, “It is clear that these and other correlations should be taken only as suggestions for further research and not as evidence of causation or as bases for preventive action.” Several studies carried out in the 1970s found correlations between fat intake and risk of breast cancer [101,102]. The association between dietary fat and risk of breast cancer appears to be weakening [103]. Goodwin and Boyd [104] reviewed a large number of studies and found that 7 of 13 international comparisons found a correlation between fat intake and breast cancer risk but only 1 of 14 case-control studies did. Hirohata et al. found no association between dietary fat intake and breast cancer in Japan [105] or Hawaii [106]. The NHANES I reported on 99 cases of breast cancer, as opposed to 5386 noncases, and found no differences in fat or fatty acid intake [107]. Willett et al. [108] studied a cohort of more than 89,000 American women whose fat intake ranged from 32% to 44% of calories and found a slight decrease in relative risk with increasing fat intake.

As in the case of breast cancer, local (case-control) studies of fat intake and colon cancer risk show minimal correlations with fat intake [109–111] whereas international studies find strong correlation [100,112]. Rogers and Longnecker [113] reviewed diet and cancer and in a summary of 24 cases of colon cancer found a small, but inconsistent, association between fat intake and risk of colon cancer. Stocks and Karn [114] in an early (1933) study of colon cancer and diet in England found dairy foods to be negatively correlated with risk. Jensen et al. [115] in studying colon cancer and diet in Finland and Denmark found saturated fat to be inversely correlated with risk. Stemmermann et al. [116] made a similar observation in Hawaii. Others [117,118] reported that risk increased with increasing intake of saturated fat. Tuyns et al. [119] in a Belgian study suggested that the dietary factor leading to increased risk was oligosaccharides and not fat. It should be evident from the fore-
going that there is no consensus with regard to the effects of dietary fat on colon cancer risk.

Is it fat or the calories it provides that affects cancer risk? In 1927 Hoffman [120] suggested that the increase in cancer incidence seen then was due to overnutrition. About 50 years later, Berg [121] made a similar suggestion, namely, that increasing risk of hormone-related tumors was due to increased caloric intake. Moreschi [122] showed in 1909 that transplanted tumors did not grow as well in underfed mice as in freely fed ones. A few years later Rous [123] demonstrated that neither spontaneous nor transplanted tumors showed optimal growth in rodents whose food intake was restricted. Lavik and Baumann [124] studied dietary effects on chemically induced skin tumors in mice. When the diet was low in both calories and fat, tumor incidence was nil. Tumor incidence in mice fed diets low in fat but high in calories was 48% higher than in those whose diet was low in calories but high in fat and only 18% lower than that in mice fed a high-fat, high-calorie diet (Table 4). Our work [125–128] showed that caloric restriction by 40% led to significantly reduced incidence of chemically induced breast or colon tumors in rats when the restricted diet contained twice as much fat. At 10% caloric restriction incidence of induced breast tumors in rats was unchanged, but tumors per tumor bearing rat were reduced by 36% and total weight of tumors by 47%. Rats fed a diet containing 26.7% corn oil but restricted by 25% exhibited lower tumor incidence, fewer tumors, and smaller tumors than rats freely fed a diet containing only 5% fat (Table 5). In humans both colon [129,130] and gastric [131] cancer have been correlated positively with caloric intake. Overweight in humans is clearly correlated with increasing cancer risk [132–134].

The effects of fat in cancer need to be stratified by fat type, fat quantity, and total caloric intake. There is too frequently a rush to judgment concerning effects of specific nutrients. Much of the recent interest in diet and cancer can be traced to the major paper by Doll and Peto [135] in which they suggested that as many as 35% of deaths due to cancer might be associated with diet. They then made the following comment: “It must be emphasized that the figure chosen (of 35% of cancers related to diet) is highly speculative and chiefly refers to dietary factors which are not yet reliably identified.” They also stated that “there is no evidence of any generalized increase (in deaths due to cancer) other than that due to tobacco.”

### Table 4

<table>
<thead>
<tr>
<th>Level of Calories</th>
<th>Tumor incidence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>0</td>
</tr>
<tr>
<td>High</td>
<td>54</td>
</tr>
<tr>
<td>Low</td>
<td>28</td>
</tr>
<tr>
<td>High</td>
<td>66</td>
</tr>
</tbody>
</table>

*Source: From Ref. 124.*
Table 5  Influence of 25% Caloric Restriction on Dimethylbenz(a)anthracene-Induced Tumors in Rats Fed High Fat Diets

<table>
<thead>
<tr>
<th>Diet</th>
<th>Incidence (%)</th>
<th>Multiplicity&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Wt (g)&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Burden&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ad libitum</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5% fat</td>
<td>65</td>
<td>1.9 ± 0.3</td>
<td>2.0 ± 0.7</td>
<td>4.2 ± 1.9</td>
</tr>
<tr>
<td>15% fat</td>
<td>85</td>
<td>3.0 ± 0.6</td>
<td>2.3 ± 0.6</td>
<td>6.6 ± 2.7</td>
</tr>
<tr>
<td>20% fat</td>
<td>80</td>
<td>4.1 ± 0.6</td>
<td>2.9 ± 0.5</td>
<td>11.8 ± 3.2</td>
</tr>
<tr>
<td>Restricted</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20% fat</td>
<td>60</td>
<td>1.9 ± 0.4</td>
<td>0.8 ± 0.2</td>
<td>1.5 ± 0.5</td>
</tr>
<tr>
<td>26.7% fat</td>
<td>30</td>
<td>1.5 ± 0.3</td>
<td>1.4 ± 1.0</td>
<td>2.3 ± 1.6</td>
</tr>
<tr>
<td>&lt;i&gt;p&lt;/i&gt;</td>
<td>&lt;0.005</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

<sup>a</sup>Tumors per tumor-bearing rat.
<sup>b</sup>Average tumor weight per rat.
<sup>c</sup>Weight of all tumors per rat.

Source: From Ref. 128.

A fatty acid that has been known for decades has recently emerged as a potent inhibitor of carcinogenesis. CLA has been shown to inhibit chemically induced fore-stomach tumors in mice when given intragastrically [136] and mammary tumors when included in the diet of rats [137]. It inhibits the growth and metastasis of human tumors when injected into immune deficient mice [138] and inhibits growth of tumor cells in vitro [139]. The effects of CLA on carcinogenesis have been reviewed recently [140]. Its mode of action is unknown at this writing.

We have accumulated reams of data relating to effects of specific macro- and micronutrients on cancer risk in humans and cancer development in experimental animals. We must now begin to examine the interactions of nutrients as they affect the major degenerative diseases—heart disease and cancer. It is also becoming evident that dietary patterns may be more indicative of risk than any particular dietary components.

ACKNOWLEDGMENT

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REFERENCES


66. R. Wood, K. Kubena, B. O’Brien, S. Tseng, and G. Martin. Effect of butter, mono- and polyunsaturated fatty acid-enriched butter, trans fatty acid margarine and zero trans...
nutrition examination survey. I. Epidemiologic follow-up study.  


118. A. J. Tuyns. A cooperative study of the habits, homelife, dietary and family histories of 450 cancer patients and an equal number of control patients.  

*Cancer Res.* 3:749 (1943).


123. M. Jain, G. M. Cook, F. G. Davis, M. G. Grace, G. R. Howe, and A. B. Miller. A case-control study of diet and colorectal cancer.  


I. INTRODUCTION

Fatty acids serve a wide variety of metabolic functions critical to all forms of life. They are a rich source of energy and carbon and well designed as a convenient unit for energy storage. However, the importance of fatty acids in human nutrition and physiology goes well beyond their role as a source of calories. Fatty acids provide the structure and hydrophobicity crucial to the maintenance of a semipermeable membrane barrier. Their structures can be modified by desaturation and elongation to produce a substantial variety of species with individual chemical and physical properties. Ester linkages to glycerides allow fatty acids to be easily exchanged for one another and allow cells to manipulate the physical properties of their membranes. Fatty acids also serve as precursors to active signal molecules such as eicosanoids, which are capable of producing potent biological effects. Evolution has produced a distinction between plants and animals in their capabilities for the metabolism of fatty acids. Higher animals are unable to synthesize all of the fatty acids required for certain tissue functions and are obligated to ingest fatty acids that are synthesized by plants. Animals have evolved a separate and distinct series of metabolic modifications of fatty acids, but are still unable to alter the original modifications inserted by plants. As a result, the membrane, signal, and storage lipids of animals vary widely according to their dietary intakes. In addition, the ability of an animal to produce a specific fatty acid relies either on an inherent mechanism for desaturation of saturated fatty acids or on the ingestion of a convertible precursor. The ingestion and/or metabolism of particular unsaturated fatty acids is necessary for a great variety
of physiological and cellular functions. Inadequate intake or defective metabolism leads to various dysfunctions due to deficiencies of these fatty acids in particular cellular locations. In addition, dietary fatty acids have been well correlated with metabolic and physiological alterations associated with heart disease and cancer [1–3]. Unsaturated fatty acids in particular play an important role in these non-energy-producing metabolic functions.

Dietary fatty acids have the singular ability among macromolecules to incorporate into tissue intact, thereby altering tissue acyl compositions. Proteins and nucleic acids, while providing energy and building blocks for metabolism, are incapable of remodeling the protein and nucleic acid compositions of tissues in their own image. Consequently, fatty acids occupy a unique and important role in human nutrition. With the recent advances in basic knowledge of plant fatty acid biosynthesis and genomics, it is possible to produce virtually any fatty acid in significant quantities. The availability of fatty acids for supplementation and the ability to engineer agricultural products provide opportunities to significantly modify the lipid content of the food supply. Unfortunately, knowledge concerning fatty acid function in physiology lags far behind the ability to modify dietary fatty acid compositions. Developing an understanding of specific fatty acids and interactions among fatty acids and how they affect individual metabolism and health will be critical for nutrition and agriculture in the next decade.

II. FATTY ACID BIOSYNTHESIS

The primary product of fatty acid synthase in both plants and animals is palmitic acid. However, many plant and animal fatty acids are longer and more unsaturated than palmitic acid; consequently, acyl modification systems are a critical component in the regulation of cellular acyl composition. Products of fatty acid synthase or dietary fatty acids may be modified by a chain elongation or the insertion of double bonds. Both elongation and desaturation reactions are organism-, tissue-, and cell-specific, allowing individual cells to maintain their compositional identities largely independent of diet. The possibilities and limitations imposed on the fatty acid composition of a cell are intimately associated with these enzymatic reactions. Figure 1 provides an overview of the most common mammalian fatty acid modifications.

Prior to discussing the biosynthesis of fatty acids, it is useful to describe the standard nomenclature for fatty acids. The systematic method of naming fatty acids provides information on acyl chain length and the degree and position of desaturation. Standard nomenclature describes a double bond occurring between the ninth and tenth carbons from the carboxyl end of a 16-carbon fatty acid as a Δ9 double bond (Fig. 2). The same bond, when viewed from the methyl end of the fatty acid, is referred to as an n-7 double bond. The usefulness of two nomenclature systems has grown out of a need to view fatty acids from multiple perspectives. Customarily, the n system is used when fatty acids are discussed with respect to nutrition, whereas the Δ designation is more useful when observing the biochemical reactions of fatty acids. In many ways, the n system simplifies investigation into the nutritional relevance of fatty acids. Animals are not capable of desaturating on the methyl side of a previously formed double bond; thus, in humans and other animals, fatty acids of a particular n designation will remain so permanently. This greatly simplifies the analysis of fatty acid metabolism in animals. In contrast, because many organisms,
including humans, are capable of acyl chain elongation, the Δ designation for a particular double bond is subject to a change in nomenclature with each elongation of the fatty acid. The Δ designation proves useful, however, when interpreting the chemical reactions associated with fatty acids. For instance, desaturases are referred to by the Δ designation because their catalytic action is consistent with the stereospecific insertion of a double bond from the carboxyl end of a fatty acid.

Omega (n)-designation

\[
\begin{array}{cccccccc}
16 & 14 & 12 & 10 & 8 & 7 & 5 & 3 & 1 \\
\end{array}
\]

Δ-designation

\[
\begin{array}{cccccccc}
2 & 4 & 6 & 8 & 11 & 13 & 15 \\
\end{array}
\]

Figure 1 A generalized scheme for human fatty acid metabolism. Gray bars represent desaturase activities, black arrows represent elongase or chain-shortening activity, and the relative abundance of each fatty acid is represented by its size.

Figure 2 The omega (n) and delta (Δ) numbering systems for palmitoleic acid. Δ-numbering starts at the carboxyl terminus carbon and n-numbering starts at the methyl terminus carbon.
A. Carbon Source and De Novo Synthesis

The biosynthesis of fatty acids is largely similar among plants and animals. Both are capable of producing fatty acids de novo from acetyl CoA via the concerted action of acetyl CoA carboxylase and fatty acid synthase. The first step in the de novo synthesis of fatty acids involves the production of malonyl CoA from acetyl CoA, a reaction catalyzed by acetyl CoA carboxylase. Acetyl CoA carboxylase carries out two partial reactions, each catalyzed at distinct sites, which first carboxylate the reaction cofactor biotin and then transfer the carboxyl group to acetyl CoA [4]. The net reaction is shown below:

\[
\text{Acetyl CoA} + \text{HCO}_3^- + \text{ATP} \rightarrow \text{malonyl CoA} + \text{ADP} + \text{P}_i
\]

In animals, this enzyme is soluble in the cytosol and appears to be regulated by a number of factors, including long chain acyl CoA, providing sensitivity to both de novo production of acyl chains and diet. The second general step in the production of fatty acids is to activate both malonyl CoA and the primary unit of condensation, acetyl CoA, by transferring the acyl groups to an acyl carrier protein (ACP). These reactions are catalyzed by malonyl CoA-ACP transacylase and acetyl CoA-ACP transacylase, respectively. The malonyl ACP complex then enters a cycle of elongation catalyzed by the soluble enzyme complex fatty acid synthase. Fatty acid synthase lengthens the acyl chain by two carbons per cycle of activity, using acetyl CoA as the condensing unit. This series of reactions culminates in the production of palmitic and stearic acids. The cycle is terminated when acyl ACP thioesterase hydrolyzes the acyl ACP thioester and releases a fatty acyl CoA [5]. Because the products of fatty acid synthase are consistently palmitic and stearic acids, acyl ACP thioesterase is likely to be specific for the hydrolysis of 16- and 18-carbon acyl ACP complexes. Fatty acid synthesis is extensively reviewed by Goodridge [4] and Wakil et al. [6].

Although the general mechanisms of fatty acid synthesis are similar, there are several specific differences between plant and animal fatty acid synthesis. Plant fatty acid synthase products are complexed in acyl ACP, whereas animals produce acyl CoA. De novo synthesis of plant fatty acids occurs in the plastid where the products of fatty acid synthase are predominantly palmitoyl and stearoyl ACP. These products are either utilized directly in the plastid as acyl ACP or translocated to the cytoplasm and converted to an acyl CoA complex [5]. The de novo synthesis of macromolecules in animals, including fatty acids, usually requires the transport of acetyl CoA into the cytoplasm, as acetyl CoA carboxylase and fatty acid synthetase are soluble cytoplasmic enzymes.

B. Desaturation

Fatty acid desaturation of nutritional importance to humans is largely similar in plants and animals. The requirements for desaturation include molecular oxygen, a reduced pyridine nucleotide, an electron transfer system, a terminal desaturase enzyme, and a fatty acyl substrate [7]. A net reaction scheme for the Δ9 desaturation of stearoyl CoA is as follows:

\[
\text{NADH} + \text{H}^+ + \text{stearoyl CoA} + \text{O}_2 \rightarrow \text{NAD}^+ + \text{oleoyl CoA} + \text{H}_2\text{O}
\]
Plant fatty acids provide a seminal source of polyunsaturated fatty acids (PUFAs) in human nutrition. This dependence on plant fatty acids is due to the fact that plants are capable of inserting double bonds into the Δ12 and Δ15 positions of fatty acids. The desaturation of fatty acids produced de novo by plants involves an initial Δ9 desaturation, followed by Δ12 and Δ15 desaturations. The insertion of double bonds on the methylene side of the previously unsaturated Δ9 bond is in opposition to animal metabolism, wherein additional double bonds are only inserted on the carboxyl side of the Δ9 double bond. This difference allows dietary fatty acids to be converted to forms not possible as a result of plant or animal metabolism alone. Plants express two pathways for the desaturation of fatty acids. The first, known as the prokaryotic pathway, is an array of desaturase activities that is present in the plastid. The second eukaryotic pathway requires the translocation of the fatty acids to the endoplasmic reticulum where desaturases act on the acyl groups. The products of the eukaryotic pathway are either utilized directly by the cell or translocated back to the plastid for further processing. Harwood [5] provides an extensive review on plant fatty acid desaturation. An interesting phenomenon of plant fatty acid biogenesis is that desaturation can utilize complex lipids as substrate. Both plastid-and endoplasmic reticulum–based desaturation introduce a double bond into fatty acids esterified in a phospholipid. In contrast, animals desaturate acyl CoA complexes.

Fatty acid desaturation in animals utilizes fatty acyl CoA thioesters as substrate. Despite multiple desaturase activities, cytochrome b5 reductase acts as an electron donor common to all terminal desaturases. The desaturases are present on the cytoplasmic face of the endoplasmic reticulum. Animals possess Δ9, Δ6, and Δ5 desaturase activity. Like desaturases in plants, each of the desaturases has preferred substrates, which can display organism- and tissue-specific differences. Because double bonds are found inserted at the Δ4 position of 22:6n-3 and 22:5n-6, it was originally assumed that animals also possessed a Δ4 desaturase. Such a desaturase has not been identified, and work by Sprecher and colleagues [8,9] has demonstrated that 22:6n-3 is produced via a Δ4 desaturase-independent pathway. Fatty acids containing a Δ4 unsaturation are in fact the product of an additional elongation, a Δ6 desaturation, and a two-carbon chain shortening that takes place in the peroxisome [9–11]. An interesting phenomenon specific to this process is the coordination of the movement of fatty acid from the endoplasmic reticulum to the peroxisome and back again for acylation into lysophospholipids. How the cell recognizes the production of 22:6n-3 within the peroxisome and spares it from further chain shortening is not understood.

The first double bond inserted into the saturated acyl CoA products of fatty acid synthase is at the Δ9 position and is catalyzed by stearoyl CoA desaturase. In contrast to desaturation in plants, further desaturation occurs only on the carboxyl side of the initial unsaturation. Interestingly, most PUFAs are of plant origin, as monounsaturated n-9 fatty acids produced de novo are not further desaturated by animals except in times of essential fatty acid deficiency. However, fatty acids previously polyunsaturated by plants are readily desaturated to form familiar PUFAs, such as arachidonic acid and docosahexaenoic acid (DHA). These highly unsaturated fatty acids are products of both plant and animal unsaturation, and they have double bonds inserted on both sides of the original Δ9 double bond.
1. \( \Delta 9 \) Desaturase

The stearoyl CoA desaturase enzyme (SCD) introduces a double bond at the \( \Delta 9 \) position of stearoyl CoA, forming oleic acid and the n-9 family of fatty acids. This enzyme also adds a double bond at the n-7 position of palmitic acid, forming palmitoleic acid and the n-7 family of fatty acids. The enzyme appears to be a fundamental gene product in the regulation of a host of cellular processes. Its structure has been resolved to 2.6 Å and various aspects of the functional domains were identified in the plant enzyme [12]. The sequences of various SCD genes were reported, and more recent work has focused on the upstream regulatory regions of the genes from both microbial and animal sources [13–15]. From this work, a host of interesting regulatory sites have been identified, implying that the SCD genes are multiply regulated in all cellular systems in which they are expressed. Primitive, single-cell organisms are known to alter SCD transcription in response to temperature, environmental shocks (pathogenic and osmotic), and substrate modification [16–19].

Not surprisingly, in addition to its many cellular actions, SCD appears to play multiple roles in higher plants and animals. Many reports have associated SCD with whole-tissue functions such as adipose accretion, lipid secretion, and tissue responses to stress. For example, plants induce SCD on thermal shock and senescence [20].

The most intriguing aspects of SCD in lipid regulation in animals are found in the apparent role of the SCD genes in lipogenesis. Much of this information was summarized by Ntambi [14]. Two highly homologous genes, described in mice, that code for SCD are termed, logically, scd1 and scd2. \( \text{scd1} \) is one of the first genes induced during adipocyte differentiation and this induction is responsive to insulin, carbohydrate, and elevated cAMP [15]. Furthermore, inhibition of SCD prevents adipocyte differentiation and, in mature cells, reduces lipogenesis in adipocytes and in hepatocytes of avian [21] and mammalian cells. Genetically obese animals exhibit greater amounts of SCD, consistent with a pivotal role in adipocyte function [22]. The striking ability of PUFA to down-regulate the enzyme activity is well described biochemically and recently was shown to be due to a substantial decrease in the stability of \( \text{scd1} \) mRNA [23]. It is not yet clear precisely what advantage such a regulatory control serves. However, various suggestions have been advanced that this could be a partial basis for reduced hepatic lipoprotein secretion during PUFA feeding and could possibly reduce adiposity in PUFA-fed animals. Thus, there is abundant evidence that unsaturated families of fatty acids interact with each other in highly complex ways.

The SCD gene is known to be sensitive to a variety of hormonal signals. For example, SCD is differentially regulated in females relative to males and its higher activity in females may be part of the spectrum of lipid metabolic changes that are a consequence of sex differences in the activity of growth hormone [24,25].

2. \( \Delta 12 \) Desaturase

The \( \Delta 12 \) desaturase of plants is responsible for the conversion of oleic acid to linoleic acid and is thus the molecular basis for the n-6 family of fatty acids. Although it is present solely in plants, its functional requirement in plants is not completely understood [26]. At the present time, the most apparent action of the enzyme activity is to improve thermal tolerance, especially of specific membrane compartments in...
plants [27]. However, the requirement by animals for the n-6 family of fatty acids is not solely related to thermal tolerance, and it is intriguing that a completely separate functionality has evolved in animals to take advantage of the n-6 family of PUFAs as precursors to more unsaturated species. The protein structure of the Δ12 desaturase at the molecular and sequence level is known, and the ability of the single gene product to affect Δ12 desaturation in transfection experiments with Saccharomyces is known [28]. Although structural studies have not reached the same level of understanding for the stearoyl CoA desaturase, similar functional themes are apparent, such as analogous required histidine residues [29].

3. Δ15 Desaturase

The Δ15 or n-3 desaturase catalyzes the conversion of linoleic acid to α-linolenic acid (as the respective acyl CoA). This enzyme activity is widely distributed in plant tissues and is the basis for the abundance of its products, the n-3 family of fatty acids, in plants and animals. Although these fatty acids are conspicuously enriched in thylakoid photosynthetic membranes, the molecular advantage that the n-3 double bond provides has not yet been determined. At the molecular level, there are two isoforms of the gene, whose products are located in either the microsomal fraction or plastid [29]. The plastid form of the enzyme appears to be associated with photosynthesis because it is induced by light [30]. The gene product may also have a role in altering membrane composition as a stress response as one of the n-3 desaturase genes is induced by wounding [31].

4. Δ6 and Δ5 Desaturases

Although ubiquitous, the Δ6 and the Δ5 desaturase enzymes have only recently been cloned from animals [32,33]. Interestingly, although the Δ6 desaturase is exceedingly rare in the plant kingdom, it had previously been cloned and functionally characterized from borage [34]. Like animal SCD, these desaturases utilize a cytochrome b₅ domain as an electron transfer system [32,33]. The expression of the Δ6 desaturase was presumed to be limited to hepatic tissue; however, Northern blot analyses now demonstrate that many tissues—including heart, kidney, lung, skeletal muscle, and, notably, brain—express the desaturase [33]. There is a broad distribution of Δ5 desaturase mRNA expression among tissues as well, with Northern analyses identifying expression in lung, skeletal muscle, placenta, kidney, and pancreas [32]. Like the Δ6 desaturase, the Δ5 desaturase is expressed most abundantly in liver, brain, and heart [32]. The abundance of Δ6 and Δ5 desaturase mRNA in tissues not typically found to have desaturase activity may indicate a posttranslational regulation of desaturase activity in these tissues. However, it is still an open question as to whether tissues such as heart and brain are capable of fatty acid desaturation. Although surprisingly few functional characterizations of the Δ5 and Δ6 desaturase genes have been reported, there is evidence that the mRNA expression of both the Δ6 and Δ5 desaturases is regulated by nutrition. Hepatic Δ6 desaturase expression was suppressed in mice fed corn oil (containing high concentrations of linoleic acid) relative to those fed triolein [33]. Although oleic acid is not a typical substrate for the Δ6 desaturase and, by contrast, linoleic acid is an excellent substrate, these results may indicate that the Δ6 desaturase is regulated by the degree of membrane unsaturation. Δ5 desaturase mRNA expression also appears to be regulated by dietary fatty acids.
Mice fed triolein or a fat-free diet exhibited marked increases in hepatic Δ5 desaturase mRNA expression relative to mice fed safflower oil or fish oil [32].

Although the metabolism of n-3 and n-6, and even n-9, fatty acids in animals is thought to utilize the same Δ6 and Δ5 enzymes, there are specific differences in the desaturation efficiencies with regard to substrate. For instance, the most abundant n-6 fatty acids in animals are clearly linoleic and arachidonic acids. In contrast, the only n-3 fatty acid present at concentrations commensurate with these n-6 fatty acids is DHA, despite the fact that most dietary n-3 fatty acid exists as linolenic acid (18:3n-3). There is also the virtual nonexistence of Δ6 and Δ5 desaturase products of endogenously produced or dietary oleic acid (18:1n-9) in animals. Clearly, the families of fatty acids are not simple competitors with each other for desaturation and there must exist a complex regulation of desaturation. The availability of clones for the Δ5 and Δ6 desaturase genes heralds a truly exciting and productive future for understanding the role of PUFA metabolism in physiology and nutrition.

5. Regulation of Desaturase Activity

The regulation of desaturase activity is complex and appears to involve a number of signals. The activities of mammalian desaturating enzyme systems are sensitive to several metabolic signals [35–39]. These various effectors cause an inhibition in the net desaturation through the Δ6 and Δ5 enzymes. Stearoyl CoA desaturase activity is substantially suppressed in the fasted state [40] and is restored by refeeding or insulin administration. In culture, scd2 gene expression was suppressed by cholesterol [41]. However, in rats, scd expression was induced by dietary cholesterol [42]. The regulation of SCD by PUFAs and cholesterol was reviewed by Ntambi [13].

As direct effects, the desaturases are strongly inhibited by their products; hence, diets rich in PUFAs tend to suppress desaturase activity. Strikingly, fasting also decreases Δ6 desaturation, and a basic protocol of fasting and refeeding accelerates essential fatty acid deficiency [43].

Endocrine signaling has been variously reported to affect desaturation [35, 43,44]. Glucagon, epinephrine, corticoids, and thyroxine all lower the activity of the Δ6 and Δ5 desaturases [43]. Diabetes in humans and in animal models is associated with lower concentrations of PUFAs, and this is paralleled by measurable decreases in the activity of the Δ6 and Δ5 desaturases. Consistently, insulin administration restores the desaturase activities in both humans and in animal models of diabetes and normalizes the content of PUFAs in membranes. As an illustration of the extent of control of these systems, insulin does not increase desaturation in normal individuals [45].

In studies designed to examine the mechanisms of modulatory effects, cAMP was implicated as causal to desaturase modification, and the effect can be mimicked by dibutyl cAMP both in vivo [46] and in vitro [47]. Interestingly, corticosteroids and thyroxine depress both Δ6 and Δ5 desaturase activities but increase the Δ9 desaturase activity [35,48].

To date there is a paucity of metabolic systems in which the apparent activity of the Δ5 and Δ6 desaturases increases over controls. Growth hormone strongly induces both the Δ6 and Δ5 enzyme activities and their metabolic products [25,49]. Growth hormone also substantially down-regulates the transcription of the stearoyl CoA desaturase of liver and adipose.
The reciprocal response of the Δ6 and Δ5 compared with that of the Δ9 desaturase is intriguing and may reflect a truly interactive regulation. For example, it is not clear if the suppression of the Δ9 desaturase by growth hormone is an effect solely of growth hormone or also of the metabolic products of the enhanced desaturation by the Δ6 and Δ5 desaturases. Arachidonic acid, the product of the Δ6 and Δ5 desaturases, is known to directly down-regulate the adipose SCD [23]. Such effects argue compellingly that the regulation of PUFA metabolism is sensitive to the products of metabolism whether formed de novo or ingested. This will likely be a focus of research in the future.

C. Elongation

The elongation of presynthesized fatty acids is critical to fatty acid metabolism in animals. C16 and C18 fatty acids are the primary products of both plant and animal fatty acid biosynthesis and, consequently, acyl chains longer than C18 must be elongated post de novo synthesis. Many PUFAs critical to the structure and physiology of animals are longer than C18; hence the need for an effective elongation system. In addition, the sequential desaturase activities involved in producing more unsaturated derivatives of n-fatty acids require intermediate elongation steps. This is not accomplished through further cycling of fatty acid synthase, but rather through independent activities located in the endoplasmic reticulum (ER) and the mitochondria. These systems are quite distinct and even use different substrates for condensation. Mitochondria and the ER add acetyl CoA and malonyl CoA as elongation substrate, respectively [50]. Although the mitochondria are very active in the production of acetyl CoA and β oxidation, the ER appears to possess the majority of elongase activity. The ER system for acyl elongation appears to prefer unsaturated fatty acids for further elongation, although this preference varies among tissues [50]. There is evidence that a third elongase activity is present in the peroxisomes and that it is related to the peroxisome proliferation response [51]. Although the majority of elongation occurs in the liver, other tissues also express activity.

D. Peroxisomal PUFA Synthesis

Whereas the majority of fatty acid modifications involve elongation and/or desaturation, some long chain PUFAs are produced by the removal of two carbons by one cycle of oxidation. Long chain unsaturated fatty acids are retroconverted in peroxi- somes to produce acyl chains with two or four fewer carbon units. The most common example of this type of acyl modification involves the synthesis of the long chain PUFA, DHA. DHA is produced by the elongation of 22:5n-3 to 24:5n-3 followed by a Δ6 desaturation and one cycle of β oxidation to produce 22:6n-3 [8,9,52]. There is also evidence that the Δ6 desaturase involved in this pathway is distinct from the Δ6 desaturase associated with typical PUFA production [53]. Several other fatty acids are produced by this pathway, although it is not known if this is the primary source of their biosynthesis. Voss et al. [54] showed that arachidonic acid can be produced via the retroconversion pathway in rats injected with 22:4n-6. Hagve and Christophersen [55] demonstrated that isolated rat liver cells were capable of retroconverting 22:4n-6 and 22:6n-3 to 20:4n-6 and 20:5n-3, respectively, suggesting that retroconversion is actively utilized for the production of PUFAs in vivo. The semitoxic erucic acid was retroconverted to oleic acid in cultured human fibroblasts [56].
Although it appears clear that the retroconversion of fatty acids via peroxisomal oxidation plays a significant role in fatty acid biosynthesis, several important questions remain concerning the regulation of the process. It is known that retroconversion takes place in the peroxisomes and not in mitochondria where the majority of cellular β oxidation occurs [9]. In addition, the same acyl CoA oxidase may be involved in all chain-shortening events [57]. A particularly intriguing aspect of the specific production of acyl chains by retroconversion is how the fatty acids are spared from further oxidation. Fatty acids with a Δ4 unsaturation are actively mobilized and returned to the ER where they are acylated into lysophospholipids [11]. Therefore, 22:6n-3 produced de novo in peroxisomes from an n-3 precursor is found acylated primarily in the phospholipids with remodeling pathways active at the ER. The distribution of 22:6n-3 in cellular phospholipids is dependent on whether 22:6n-3 is consumed intact or as a metabolic precursor [58]. A lack of peroxisomal retroconversion activity has been associated with several pathologies, including Zellweger’s syndrome [9,59]. The accumulation of very long chain fatty acids (C24–26) is a hallmark of Zellweger’s syndrome, suggesting that peroxisomal oxidation is both a normal and critical component of fatty acid metabolism.

III. PUFA METABOLISM IN MEMBRANES

Knowledge of tissue or even cellular fatty acid composition is not sufficient to predict the effects of PUFAs in cell physiology. The realization that fatty acid location plays an important part in the use and function of specific fatty acids has been a major advancement in lipid metabolism. Unsaturated fatty acids, as well as their saturated counterparts, are esterified in phospholipids of cell membranes. There are a variety of phospholipid types, and each type has its own unique compositional fatty acid identity. In addition to phospholipid identities, individual cell membranes have unique phospholipid and fatty acid compositions. Although there are myriad data on the fatty acid composition of plant and animal tissues, these fatty acid profiles are subject to important changes brought about by diet, disease state, or a variety of other factors. As a result of the variety of phospholipid pools in which fatty acids can be esterified, very slight changes in total cell fatty acid content can have significant effects on cell function. It is quite possible that lipids have been investigated and rejected as causal agents of a number of pathologies on the basis of largely unchanged fatty acid compositions. It is now clear that fatty acids exert their effects from specific locations and that their physiological effects are mediated in part by the movement of PUFAs into important phospholipid pools. The movement of fatty acids into phospholipid pools is catalyzed by fatty acid carrier proteins, and the acyl compositions of membranes and specific phospholipid pools are mediated by the specificities of these enzymes.

A. Remodeling

Dietary fatty acids can exert significant effects on the fatty acid compositions of phospholipid membranes. These effects must be attributed either to the synthesis of new phospholipids de novo or to activities that change the composition of preexisting phospholipids. Quite commonly, it is the latter of the two mechanisms. Phospholipid fatty acid remodeling is both an important pathway for the incorporation of dietary
fatty acids and a dynamic system for membrane property homeostasis. A generalized pathway for the remodeling of a phospholipid (Lands pathway) [60] involves (1) removal of the acyl group from the phospholipid via a lipase activity; (2) conversion of the fatty acid to an acyl CoA thioester; (3) a modification event, such as elongation or desaturation; and (4) reesterification of the acyl group or the insertion of a new acyl unit onto the lyso phospholipid (Fig. 3). The first step in the Lands pathway requires the hydrolysis of the acyl chain from the phospholipid and is catalyzed by the phospholipase A family of enzymes. Phospholipases have specificities for both the phospholipid position and acyl chains, and these specificities contribute to the regulation of membrane remodeling. Phospholipases are well reviewed by Waite [61]. Once a phospholipase has acted on an acyl chain, the fatty acid must then be activated to an acyl CoA thioester. This is accomplished via the action of acyl CoA ligases. There are several distinct acyl CoA ligase activities, each with its own cellular locations and acyl preferences [62]. The activated acyl CoA complex can be converted by desaturation or elongation (see above) and reacylated into a lyso phospholipid. The acyl transferase enzymes catalyze the esterification of acyl chains from acyl CoA thioesters into lyso phospholipids. Two types of acyl transferase activities are relevant to membrane remodeling, and not surprisingly, they catalyze the insertion of a fatty acid into either the sn-1 or sn-2 position of phospholipids. The first type of enzyme, typified by glycerol-3-phosphate acyl transferase, is capable of inserting an acyl chain into the sn-1 position of glycerol-3-phosphosphate and has a general preference for saturated acyl CoA [63,64]. In fact, glycerol-3-phosphate acyl transferase is a number of distinct enzymes capable of carrying out the same reaction at different locations in the cell, and with varying acyl CoA preferences [62,63]. The primary

Figure 3 A generalized scheme for the fatty acid remodeling of phospholipids (the Lands pathway) [60].
Acyltransferase responsible for esterifying acyl chains into the sn-2 position is lysophosphatidate acyltransferase [64]. Lysophosphatidate acyltransferase has a higher activity with unsaturated acyl CoA thioesters and is responsible for the virtual absence of saturated fatty acids at the sn-2 position in membrane phospholipids. As might be expected, the lysophosphatidate acyltransferase has a higher activity than the glycerol-3-phosphate acyltransferases [64], as the majority of the fatty acid remodeling in membranes occurs with unsaturated fatty acids. The combined preferences and activities of the acyltransferases are largely responsible for the positional and site-specific acyl compositions observed in phospholipid membranes.

1. Acyl-Specific Incorporation

Several phospholipid species are noteworthy for their high degree of incorporation of specific fatty acids. The ether-linked phospholipids, for example, are enriched at the sn-2 position by a CoA-independent transacylase relatively specific for arachidonic acid [65,66]. The most salient example of acyl-specific incorporation, however, can be found in the mitochondrial diphospholipid cardiolipin. Cardiolipin is characteristically enriched with linoleic acid to as much as 85% of its acyl species in vivo [67]. Cardiolipin is also selectively enriched in DHA to as much as 50% of its total fatty acid content [58,67,68]. The implications of the high degree of unsaturation of cardiolipin may involve its function as the membrane solvent for the electron transport enzymes. Other phospholipids incorporate specific fatty acids to varying degrees. Spector and Yorek [69] treated Y79 retinoblastoma cells with arachidonic acid, DHA, and oleic acid to determine the relative affinity of various phospholipid classes for unsaturated fatty acids. Phosphatidylcholine, phosphatidylethanolamine, and phosphatidylserine could all be enriched with the monounsaturated fatty acid oleic acid. Only phosphatidylethanolamine and phosphatidylinositol were substantially enriched in arachidonic acid, whereas phosphatidylcholine, phosphatidylethanolamine, and phosphatidylserine all incorporated docosahexaenoic acid. The differences in acyl incorporation indicate that the synthesis and remodeling mechanisms described above have different activities depending on both the fatty acid and the phospholipid substrates.

IV. DIETARY SOURCES OF UNSATURATED FATTY ACIDS

A. Microbes

Most bacteria are fully capable of synthesizing all of the fatty acids required for their normal growth and reproduction. The fatty acid synthetase enzyme complex responsible for this activity adds acetate units to a final chain length of 16–18 carbons. These can be further desaturated (normally once to a monounsaturated fatty acid) after which the saturated and unsaturated fatty acids are esterified to yield membrane phospholipids. Bacteria in general do not store energy as fats; hence, triacylglycerols are not abundant forms of lipid in bacteria. They are thus not a quantitatively important fat source in foods. The unique metabolism of bacteria, especially the production of branched chain and odd-numbered fatty acids, can occasionally generate measurable quantities of unusual fatty acids in, for example, bovine milk, due to incorporation of the products of rumen fermentation. In the
production of commercially viable oil sources, wax esters and eicosapentaenoic acid have been produced at a commercially relevant scale in bacteria [70,71].

Microbes as a broad class of single-cell (or relatively undifferentiated collections of cells) organisms have the obligate biosynthetic capability of generating fatty acids necessary for membrane synthesis and other processes. Historically these have not been considered an important source of fat in the diet, although it is recognized that these organisms add small quantities to certain foods, whose presence may be an important contribution, e.g., for flavor. Nevertheless, even for microbes that produce large quantities of storage triacylglycerides, the economics of growing and obtaining the oils has been noncompetitive with the traditional sources of edible oils. However, recently this area has seen a considerable resurgence in interest both academically and in commercial application [72]. This change is largely due to the improved efficiency and capabilities of large-scale microbial fermentation, to the identification of therapeutically useful edible oils, and to the capability of microbes to produce unusual fatty acids or unusual concentrations of fatty acids and glycerides [73]. Fatty acids that have raised the ante, as it were, for edible oils include dihomo-γ-linoleic acid, eicosapentaenoic acid, and DHA, due to their ability to alter arachidonic acid metabolism and hence thrombosis, inflammation, cancer, and autoimmune diseases; DHA for inclusion in infant formulas; nervonic acid for its potential in treating neuropathies; long chain monounsaturated fatty acids for adrenoleukodystrophy; and stearcucic acid as a possible treatment for bowel cancer [74,75]. Several factors mitigate in favor of microbial production for high-value lipids. The greater potential for aggressive recombinant approaches to manipulate microbial lipid metabolism to obtain novel fatty acids is likely to increase the growth of this cottage industry for fatty acid production. Higher plants and animals are also somewhat limited in the glyceride forms that they will produce. For example, most plants do not place a saturated fatty acid in the sn-2 position of a triglyceride. Similarly, fish tend to place virtually all of the long chain n-3 PUFAs in the sn-2 position. This both limits the total range of glycerides available using these plants and animals as sources of lipids and imposes structural effects on digestion and absorption of the fatty acids in nutritional applications. These limitations are both less well defined and more mutable in microbial fermentation applications. This area, though coming under intense regulatory scrutiny, may reach a significant segment of the food industry, at least in the short term [76,77].

Single-cell eukaryotes have, as a class, a remarkably wide variety of lipid metabolic capabilities. For example, some yeast produce only a single desaturase, the stearoyl or Δ9 enzyme, and neither produces or requires PUFA for growth. At the other end of the spectrum, some fungi and algae can produce very high amounts of arachidonic, dihomo-γ-linolenic, eicosapentaenoic, and DHA [78]. An additional and synthetically useful attribute of these organisms is their ability to take up fatty acids from the medium and either to incorporate them into triacylglycerides and phospholipids (even with unusual stereospecificity) or to further metabolize them prior to esterification [78]. These various properties were known previously but were not thought to warrant commercialization. This is changing. Already, microbial lipid sources are proving to be a cost-effective feedstock for shrimp and fish aquaculture. The ability of the microbial feedstock to also elaborate valuable pigments and antioxidants is used to advantage, so this entire technology and its biotechnological elaborations are likely to increase in impact in the future.
B. Agricultural Products

The extended metabolism of 18-carbon PUFAs to longer chain, more unsaturated fatty acids [frequently referred to as highly unsaturated fatty acids (HUFAs)] in animals means that even though animals and plants may contribute similar families of PUFAs to the diet, the precise form of these fatty acids will differ. As a result, an important consequence of consuming animal in contrast to vegetarian foods is that in the latter, linoleic and linolenic acids are the fatty acids ingested from the n-6 and n-3 families, whereas in animal foods, their metabolic products, preformed arachidonic, eicosapentaenoic, and DHA, are also ingested. It is now clear that these are significant nutritional, biochemical, and physiological differences.

Furthermore, whereas animal sources of fat are often grouped as similar, avian, aquatic, and ruminant or nonruminant mammalian storage lipids are very different in the quantity of depot triacylglycerols, their distribution and their fatty acids, as well as their composition and arrangement on the glycerol. The final content of fatty acids in storage triacylglycerols is the result of diet, metabolism, and de novo synthesis. In this respect, each of the major animal fat sources differs in important ways, which tends to distinguish each as a fat-rich commodity. These differences have important effects on the texture, flavor, and caloric density of the muscles as consumed directly [79–81] and also on processed foods prepared from them [82].

Although the differences among species in the quantity and distribution of fat are associated with the particular commodities, they are not necessarily all innate to them. These differences also reflect the historical development of the particular muscle food as a commodity. Even among ruminants, the fat content of modern beef muscle is higher and more saturated than that of comparable wild ruminant muscle [83]. Breeding and feeding practices allow for the production of meat at a specific fat concentration [84]. If different properties were perceived to be beneficial, the fat content could arguably be altered to various extents accordingly. Thus, when examining the content of storage fat in muscle tissue that is used as food, one is looking at a rather narrow window of a wide range of possibilities. As commodity needs become more defined and the fat functionality better understood, the means to arrive at these targets will need to be explored.

In addition to the differences in total quantity of fat and its tissue distribution, the composition of storage triacylglycerols in animal species differs as well. Red meats tend to be relatively higher in saturated fatty acids and lower in PUFAs than poultry or fish. Poultry and fish differ significantly in the chain length of monounsaturated fatty acids and in the content of n-3 PUFAs.

Once again, a consistent observation of the lipid content in different animal tissues is the variability within species. Even within ruminant animals in which the dietary PUFAs are largely hydrogenated by rumen flora, there is a significant range of composition. Among monogastric animals, the variability in fat composition within species due to muscle type, diet, environment, and age is typically greater than the differences noted among species [81].

An important question becomes, what unique properties of the metabolism of the three animal types lead to the observed or apparent differences in lipid composition and behavior? In all animals, the storage triacylglycerols both in adipose and individual muscle cells can be assembled from both dietary fatty acids and fatty
acids synthesized de novo, primarily either in liver (in chickens and fish) or in adipose tissue (in pigs) [85,86]. In general, de novo synthesis of saturated fats is decreased by dietary fats [81]. Therefore, fats from the diet constitute the greatest source of variation in the composition of storage fats. Within this framework, metabolic control can be seen. For example, short and medium chain fatty acids are not incorporated into storage lipids of most animals. These pass into the liver where they are either elongated or oxidized for fuel [87]. Although monounsaturated and linoleic acid, 18:2, are readily incorporated, in most animals long chain (greater than C18), highly unsaturated fatty acids are not esterified into triacylglycerols [85]. However, fish will accumulate HUFAs, notably the n-3 PUFAs 20:5 and 22:6, but only if they or their precursors are present in the diet and only at low water temperatures [88]. Fish actually require n-3 PUFAs in their diets but are unable to synthesize them [89,90]. Alternatively, very high concentrations (>50%) of saturated fats are not found in storage lipids due to the well-regulated activities of the Δ9 desaturase that produces oleic acid from stearic acid [85]. In ruminant animals, the rumen microorganisms hydrogenate unsaturated fatty acids in the diet, which has an overriding influence on the composition of the storage fats [91]. However, when PUFAs such as 18:2 are protected from ruminant microorganisms, they accumulate in storage lipids in beef comparably to accumulation in nonruminants [50]. Finally, mammals absorb fat into the lymph, whereas fish and poultry absorb fat directly into the portal vein. As a result, adipose tissue can access incoming fatty acids directly in mammals, but fat passes by liver first in avians and fish. Thus, there is considerably more hepatic metabolism of ingested fatty acids in avian and fish tissues.

C. Effect of Agriculture on the Composition of the Food Supply

Fatty acids occupy a unique position in nutrition in that they have the ability to survive digestion intact, enabling them to replace the fatty acid content of the consumer. It is reasonable then to expect that the lipid composition and, correspondingly, the physiology of individuals who consume particular fats and oils to be reflective of the fatty acid composition of their diet. Interestingly, it has been postulated that the dietary PUFA composition of an average human diet has changed markedly with modern advances in agriculture [92–94]. Wild foods are typically much higher in n-3 PUFAs than crops successfully developed by agriculture. There is a variety of evidence to suggest that the changing ratio of n-3 to n-6 fatty acids has affected human physiology adversely and that humans may have developed major classes of pathologies as a result of this change. The lower rates of coronary heart disease and cancer in populations consuming a higher n-3 to n-6 PUFA ratio are well documented [95–97]. Despite mounting evidence that human populations would benefit from an increased consumption of n-3 fatty acids, it is unlikely that this change will occur in the near future. The primary reason for this is the agricultural success of crops rich in n-6 fatty acids. The n-6 fatty acids are most typically found in seed crops, which are not only consumed directly but are also used in animal feed. In addition, n-6-rich crops are generally more stable than n-3-rich crops, leading to their preferential cultivation and use as food ingredients. Thus, the increases in coronary artery disease, cancer, and autoimmunity may be a direct consequence of the advance of modern agriculture.
V. NUTRITIONAL EFFECTS OF UNSATURATED FATTY ACIDS

The field of PUFA biochemistry has only begun to develop convincing molecular models for the effects of fatty acids on physiology. Having lagged somewhat behind, lipid biochemistry is now poised to develop in the same way that protein and nucleic acid biochemistry has over the last 20 years. There is considerable information known about fatty acid synthesis as well as a massive collection of data on the fatty acid composition of foods. It has recently even become feasible to modify the fatty acyl content of foods through genetic manipulation. Yet there are very few data on how and why particular fatty acids modulate physiology. An understanding of the molecular basis of fatty acid nutrition will be important for the design of diets appropriate to the individual. PUFAs are thought to exert their physiologic effects through a variety of mechanisms, ranging from acting as precursors for signal molecule formation to modulating membrane structure. The remainder of this chapter will review what is known about the non-energy-producing functions of PUFA and how individual fatty acids modulate these functions.

A. Role of PUFA in Cell Physiology

Whereas all fatty acids contribute hydrophobicity to membranes, unsaturated fatty acids provide several unique functionalities. Unsaturated fatty acids and particularly PUFA can form a vast array of chemical structures, each of which has unique physiochemical properties. Cells can use fatty acids to modulate their membrane properties and the activities of membrane-associated enzymes, and for the production of potent signal molecules. Knowledge of the effect of fatty acids on membrane properties has been impeded by the fact that it is difficult to greatly modulate membrane fatty acid composition. Although the difficulty associated with modifying cell membrane composition suggests that membrane homeostasis is critical to cells, it makes investigation on the effects of individual fatty acids difficult. The true successes in this area have come from the discovery of small but highly active phospholipid pools and the enzymes that maintain them. The regulation of the ether-linked arachidonate-containing phospholipid pools is the best described example of how small changes in membrane compositions can have significant effects on cell physiology [98]. Future research in this area will likely have to focus on developing controllable models in which the contributions of specific fatty acids are determined.

1. Unsaturated Fatty Acids and Membrane Structure

A critical feature of the production of unsaturated fatty acids by eukaryotes is that the carbon–carbon double bonds exist in the cis configuration. Pi-bonded carbon–carbon double bonds may exist in two conformations, cis and trans (see Fig. 4), yet nature has carefully preserved the production of the cis isomer to the virtual exclusion of trans isomers. The reason for this is best explained by viewing fatty acids as important structural components of cells. Brenner [99] and Cook [50] reviewed the key physiochemical features of double bonds in fatty acids. Most important among the changes imparted to a fatty acid by a cis double bond is the rigid “kink” or bend in the acyl chain.

Membranes are largely held together by London–van der Waals forces between adjacent fatty acyl chains [50]. Because these interaction forces are significantly diminished with even a slight increase in the distance between acyl chains, fatty acid
packing plays a key role in membrane structure. The placement of unsaturated fatty acids into membranes is confined, with few exceptions, to the sn-2 position of phospholipids; as a result, the unsaturated fatty acid content of membranes rarely if ever exceeds 50 mol % unsaturated acyl chains. An advantage of including both an unsaturated and a saturated fatty acid on the same phospholipid molecule is that the two types of acyl chains cannot demix in the membrane. This ensures that slight depressions in the energy of association, such as those achieved between the spontaneously formed sphingomyelin-cholesterol in raft complexes, are not achieved with individual fatty acid types. The association between adjacent phospholipid molecules is entirely dependent on the van der Waals forces that attract their fatty acid components. These effects can be attributed to two primary structural properties of the fatty acid: (1) the length of the acyl chain and (2) the degree of unsaturation. These two structural elements can significantly affect both the relative volume the fatty acid occupies and the distance between a fatty acid and its neighboring acyl chain. The degree of unsaturation is particularly important for membrane properties because van der Waals forces are acutely sensitive to the distance between the interacting acyl chains [50].

Estimates of the spatial widths of fatty acids show significant differences between cis and trans double bonds. Whereas stearic acid is estimated to be 0.25 nm in diameter, Δ9-cis-octadecanoic acid (oleic) and Δ9-trans-octadecanoic acids have spatial widths of 0.72 and 0.31 nm, respectively [50]. Additionally, due to the fact that the acyl chain continues on the same side of the double bond, the cis configuration imparts a 30° bend in the fatty acid that is not relievable through any rotation.
of the single-bonded carbon atoms. In contrast, the structure of a trans fatty acid is relatively unaffected by the double bond [50]. The cis unsaturated bond thus interrupts a succession of London–van der Waals forces between membrane fatty acids by increasing the distance between adjacent fatty acids and lowers the membrane crystallization temperature. The more unsaturated the fatty acid, the less it is able to rotate around its carbon–carbon bonds and, consequently, the more it influences membrane acyl packing. Some calculations suggest that an increase in the number of double bonds provides diminishing returns in terms of the fatty acid’s influence on van der Waals forces [99]. It is clear that acyl unsaturation can affect van der Waals interactions through either changes in the length of association or by increasing the distance of acyl separation. In addition, the position of a double bond in a fatty acid also plays a role in the modification of membrane structure. Double bonds at the methyl end of acyl chains are not particularly effective at modulating membrane bulk properties due to the fact that they induce a lesser degree of acyl separation. Interestingly, double bonds are typically first inserted at the Δ9 position, and fatty acids with double bonds present only near the methyl end of the chain are conspicuously absent from nature.

Cells take advantage of the large variety of fatty acid structures to maintain consistent bulk membrane properties in the face of changing temperature and pressure. Clearly, then, PUFAs are critical to cell function if viewed only from the perspective of membrane structure. In light of this fact, it is interesting that animals are largely incapable of producing PUFAs de novo.

2. Alterations in Membrane-Associated Enzyme Activity

Many membrane-associated enzymes are responsive to their fatty acid environment. There is considerable evidence that membrane unsaturated fatty acid content can modify the structure and therefore the functionality of membrane enzymes, and these effects are often cited when dietary fatty acids modulate physiologic functions. The direct action of membrane composition on membrane-associated enzyme activity is often difficult to establish. However, an interesting line of experimentation that involves measuring membrane-associated activities as a function of temperature has provided compelling evidence that the degree of membrane unsaturation is critical to cell function. Membrane-associated enzyme activity [100], molecular transport [101], and the insertion of proteins into membranes [102] all have maximal activity at temperatures just above the transition temperature of the membrane. Although clearly this does not indicate that cells use the changes in membrane unsaturation to modulate enzyme activities, it does provide strong evidence that diet-induced changes in membrane composition could play an important role in the modification of cell physiology.

For all of the difficulties in assigning a causal mechanism to changes in membrane-associated enzyme activities, it is worth noting that the effects themselves are largely incontrovertible. Dietary oils and fats have significant effects on many metabolic activities, including the activity of membrane-associated enzymes. An increasing body of literature has amassed concerning the modulation of membrane-bound enzyme activities in both cell and reconstituted systems. Brenner [99] provides an extensive review of the effects of unsaturated fatty acids enzyme kinetics.
VI. SYNTHESIS AND ABUNDANCE OF PUFA

Despite the considerable variation that is possible in PUFA structure, there are only about 20 unsaturated fatty acids of nutritional importance to humans. These consist primarily of monounsaturated and methylene-interrupted PUFAs in the cis configuration. Virtually all of the unsaturated fatty acids consumed in normal diets are members of the n-3, n-6, n-7, or n-9 families of fatty acids. Non-methylene-interrupted and trans-configured fatty acids were historically consumed in very small quantities, but the advent of modern food production may have enriched these fatty acids in the food supply. The primary unsaturated fatty acids in human nutrition are reviewed below.

A. n-7 Fatty Acids

1. Palmitoleic Acid (16:1n-7)

Palmitoleic acid is a minor component of both animal and vegetable lipids. Fish oil is particularly enriched in palmitoleic acid, and some seed oils also represent a significant source of the fatty acid. Palmitoleic acid is produced de novo by plants and animals by the Δ9 desaturation of palmitic acid.

2. Vaccenic (18:1n-7)

Vaccenic acid is a major product of bacterial fatty acid synthesis and is also present at lower concentrations in plant and animal lipids. Vaccenic acid is produced by the elongation of palmitoleic acid. The true content of vaccenic acid in the diet may be underestimated due to difficulty in separating it from its n-9 isomer. In animals, vaccenic acid appears to be concentrated in the mitochondrial lipid cardiolipin [103].

B. n-9 Fatty Acids

1. Oleic Acid (18:1n-9)

Oleic acid is a Δ9 desaturase product of stearic acid and is produced de novo in plants, animals, and bacteria. Oleic acid is the most common unsaturated fatty acid and is the precursor for the production of most other PUFAs. Plants produce both n-3 and n-6 PUFAs from oleic acid, and animals can elongate and desaturate oleic acid into a variety of n-9 fatty acids. Olive oil is a particularly rich dietary source, and most foods, especially nuts and butter, are rich in oleic acid.

2. Erucic Acid (22:1n-9)

Erucic acid is a long chain monounsaturated fatty acid found in plants, particularly in rapeseeds. It is an elongation product of oleic acid, and is an uncharacteristically long chain unsaturated fatty acid for plants. Mildly toxic, erucic acid has been bred out of rapeseeds used for food oil production. In animals, dietary erucic acid can be retroconverted to form oleic acid via peroxisomal oxidation.

3. Mead Acid (20:3n-9)

Mead acid is a hallmark of essential fatty acid deficiency and has the distinction of being the only major PUFA produced de novo by animals. In the absence of dietary n-6 and n-3 fatty acids, the Δ6 desaturase converts oleic acid to 18:2n-9, which is
further elongated and \(\Delta 5\)-desaturated to form mead acid [50]. It has been speculated that mead acid compensates for the loss of n-3 and n-6 PUFAs by increasing the unsaturation of animal cell membranes.

4. Other n-9 Fatty Acids

The family of n-9 fatty acids is derived exclusively from the production of oleic acid but can be converted by elongation, desaturation, \(\beta\) oxidation, etc. Other rare but naturally occurring n-9 fatty acids include 18:2, 20:1, and 22:3 [104].

C. n-6 Fatty Acids

1. Linoleic Acid (18:2n-6)

Linoleic acid, along with \(\alpha\)-linolenic acid, is a primary product of plant PUFA synthesis. Linoleic acid is produced de novo by plants and in particular is enriched in seed oils. Although nature produces linoleic acid at concentrations fairly equitable with those of \(\alpha\)-linolenic acid, modern agriculture has greatly enriched linoleic acid in the food supply. Although animals are incapable of producing linoleic acid, livestock are fed diets particularly rich in this fatty acid, and thus humans acquire a large portion of their linoleic acid from meats. Linoleic acid serves as a precursor for the production of the essential fatty acid arachidonic acid, as well as other n-6 acyl species.

2. \(\gamma\)-Linolenic Acid (18:3n-6)

\(\gamma\)-Linolenic acid (GLA) is produced in animals and lower plants by the \(\Delta 6\) desaturation of linoleic acid. Natural sources include evening primrose oil, borage oil, and black current oil, and minute amounts can be found in animal tissue [105]. In animals, dietary linoleic acid is desaturated by the \(\Delta 6\) desaturase to produce \(\gamma\)-linolenic acid as an intermediate in the production of arachidonic acid. Interestingly, dietary \(\gamma\)-linolenic acid is accumulated in animal tissue largely as its direct elongation product 20:3n-6, and not substantially converted to arachidonic acid. There has been a great deal of recent interest in dietary \(\gamma\)-linolenic acid for its antagonistic action on arachidonic acid metabolism.

3. Dihomo-\(\gamma\)-Linolenic Acid (20:3n-6)

The elongation product of linoleic acid, dihomo-\(\gamma\)-linolenic acid (DGLA), is a minor component of animal phospholipids. DGLA serves as a precursor to the formation of the essential fatty acid arachidonic acid as well as for the prostaglandin \(G_\text{I}\) series. Dietary DGLA does not appear to be rapidly converted to arachidonic acid, and because prostaglandins of the \(G_\text{I}\) series have anti-inflammatory properties, DGLA has received attention as a potential therapeutic agent.

4. Arachidonic Acid (20:4n-6)

Arachidonic acid is the product of desaturation and elongation of linoleic acid in animals. Arachidonic acid is also produced in quantity in marine algae. Dietary linoleic acid is converted to arachidonic acid in animals by the concerted activity of the \(\Delta 6\) desaturase, a microsomal elongase and the \(\Delta 5\) desaturase. Arachidonic acid is referred to as an essential fatty acid for its action as the precursor for the production
of eicosanoids. It is present in all tissues and is particularly enriched in phosphati-
dylcholine, and ether-linked phospholipid membrane pools.

5. Docosatetraenoic Acid (22:4n-6)

Docosatetraenoic acid is the direct elongation product of arachidonic acid and is present in minimal amounts in animal tissues. Docosatetraenoic acid is a substrate for peroxisomal retroconversion resulting in the formation of arachidonic acid [55].

6. Other n-6 Fatty Acids

The family of n-6 fatty acids is derived exclusively from the production of linoleic acid but can be converted by elongation, desaturation, β oxidation, and so forth. Other rare but naturally occurring n-6 fatty acids include 16:2, 20:2, 22:2, 22:3, 24:2, 25:2, 26:2, and 30:4 [104].

D. n-3 Fatty Acids

1. α-Linolenic Acid (18:3n-3)

α-Linolenic acid is produced de novo by the Δ12 and Δ15 desaturation of oleic acid in plants. Along with linoleic acid, α-linolenic acid constitutes one of the two primary PUFA products of plant fatty acid biosynthesis. It is primarily present in the leaves of plants but is also a minor component of seed oils. α-Linolenic acid serves as the metabolic precursor for the production of n-3 fatty acids in animals. The success of agricultural seed oils has caused a significant shift in the natural balance of linoleic and linolenic acids, and over the last 100 years the average dietary content of α-linolenic acid has declined significantly [94].

2. Eicosapentaenoic Acid (20:5n-3)

Eicosapentaenoic acid (EPA) is produced de novo by marine algae and in animals by the desaturation/elongation of α-linolenic acid. EPA is the primary fatty acid of fish oil (approximately 25–20% by weight) although it is not produced de novo by fish. It has also been reported that significant EPA production can occur in animals by the β-oxidation chain shortening of DHA [55]. EPA has been investigated extensively for its action as a competitive inhibitor of arachidonic acid metabolism. Although eicosanoids can be produced from EPA, they appear to have either no activity or an activity that opposes arachidonic acid–derived eicosanoids.

3. Docosapentaenoic Acid (22:5n-3)

Docosapentaenoic acid is the elongation product of EPA and is present in most marine lipids. Docosapentaenoic acid can be converted to DHA via a three-step process involving a unique Δ6 desaturation in animals (see above) [8].

4. Docosahexaenoic Acid (22:6n-3)

Docosahexaenoic acid (DHA) is produced de novo by marine algae and is a primary component of fish oil (approximately 8–20% by weight). The production of DHA in animals from linolenic acid occurs via the desaturation/elongation of α-linolenic acid to 24:5n-3. This very long chain unsaturated fatty acid is desaturated by a Δ6 desaturase (possibly a unique Δ6 desaturase enzyme) and the resulting fatty acid undergoes one cycle of β oxidation to form DHA [8,9]. Animals appear to have a
requirement for DHA for neural function and they rely on its production from n-3 precursors by elongation/desaturation cycles or through ingestion of the intact acid [106]. Although the exact role DHA plays in animal physiology is not understood, the great care with which the fatty acid is preserved in certain tissues implies that it may be an essential component of certain cells. Brain and retinal tissues are particularly enriched in DHA.

5. Other n-3 Fatty Acids

The family of n-3 fatty acids is derived from α-linolenic acid but can be modified by chain elongation, desaturation, β oxidation, etc. Naturally occurring but rare n-3 fatty acids include 16:3, 16:4, 18:4, 18:5, 20:2, 20:3, 20:4, 21:5, 22:3, 24:3, 24:4, 24:5, 24:6, 26:5, 26:6, 28:7, and 30:5 [104].

E. Unusual and Non-Methylene-Interrupted Fatty Acids

The vast majority of PUFAs contain multiple double bonds in a 1,4-pentadiene structure in which a single methylene carbon is positioned between the two double bonds. More double bonds are added as a direct result of the positional selectivity of the subsequent desaturase enzymes. Hence, from fatty acids containing two double bonds (linoleic acid 9,12 18:2) to those containing six double bonds (DHA; 4,7,10, 13,16,19), all exhibit methylene interruption over the entire length of the molecule. However, there are certain naturally occurring fatty acids in which single double bonds exist at a distance. These are the non-methylene-interrupted fatty acids (NMIFAs). Various NMIFAs have been described, including allenic, conjugated, allylic, enoic, acetylenic, cyclic, branched, hydroxylated, iso and anteiso fatty acids [104]. The most common NMIFAs are fatty acids in which one of the double bonds is ostensibly missing from the middle of a double-bond system. Conifers were shown to contain up to 20% by weight in their seeds of the NMIFA 5,11,14-eicosatrienoic acid and 5,11,14,17-eicosatetraenoic acid [107]. In most reported studies, the basis of the synthesis of these fatty acids is through an active elongase enzyme that elongates 18- to 20-carbon fatty acids, bypassing the Δ6 desaturase. Indeed, it has been argued that the methylene-interrupted structure is not due to explicit enzyme specificity but rather the predisposition of substrate fatty acids [108]. Desaturation at the Δ5 position of the elongated fatty acid produces a 20-carbon fatty acid without the Δ8 double bond. This aggressive elongation has been argued to be the basis for the occurrence of small quantities of 5,11,14-eicosatrienoic acid in animal tissues. While most unusual fatty acids are not readily esterified into membrane phospholipids, for certain structures this is not true. 5,11,14-Eicosatrienoic acid, which is the structural analog to arachidonic acid with the absence of the Δ8 double bond, is esterified into several membranes, and even shows preference for incorporation into specific phospholipids including phosphatidylinositol [107,109]. Because the absence of the Δ8 double bond makes it impossible to synthesize prostaglandins or leukotrienes from NMIFA, replacement of arachidonic acid with NMIFA in phospholipids would be predicted to have substantial effects on eicosanoid cellular signaling proportional to the extent of displacement of arachidonic acid. This was shown to be true for isolated tissues [107]. Furthermore, in an animal model of genetic autoimmunity, animals fed diets containing these fatty acids exhibited a net reduced severity of disease consistent with a selective decrease in eicosanoid signaling [110]. Other examples of un-
usual NMIFA structures with potential biological actions include conjugated linoleic acid that has exhibited potent anticancer properties in animal models of carcinogenesis [111,112]. The true therapeutic value, as well as the potential toxicities, of these fatty acids will await the development of commercial sources in food grade quantities for larger studies. Nevertheless, the potential for modifying the biosynthetic capabilities of crop plants is already being exploited for many nutritional and functional targets, and the possibility of producing fatty acids with unusual structures is likely to be limited only by the documented value of producing them.

VII. SUMMARY

The fatty acid nomenclature system based on n designation was developed to describe fatty acids in terms of their nutritional functions. Whereas this approach is of some use, it may lead to confusion over the true basis by which fatty acids modulate physiology. There is mounting evidence that each fatty acid has its own role in nutrition that is not dictated by its n designation. Chemically similar fatty acids often have widely ranging functionalities. This phenomenon is exemplified by the antagonistic relationship between arachidonic acid and EPA metabolism despite the fact that the fatty acids differ only by the additional double bond in EPA. The fundamental relation between members of the same n family is one of interconvertibility. Families are grouped only by their potential to be converted to longer chain members of the same n designation or, in a more practical sense, by their ability to act as precursors for the production of a particular fatty acid of interest. The enzymatic activities required for the conversion of fatty acids to longer, more unsaturated chain members of the same n family are redundant. Consequently, if cells were not capable of recognizing fatty acids based on their n designation and responding by modulating desaturase and elongase activities appropriately, animals would be completely at the mercy of their diets.

The regulation of acyl content appears to be critical to the function of a cell, as mammalian cells expend a substantial amount of energy in maintaining distinct and heterogeneous membrane fatty acid compositions. Cells are capable of preserving these compositional identities even when confronted with phospholipid diffusion and vesicular transport, suggesting that acyl composition is tightly regulated. In addition to overall membrane acyl content, phospholipid acyl composition is rigorously maintained in a positionally specific manner. The complex framework of enzymatic activities that upholds these compositions makes large-scale changes in membrane composition rare and, as a result, the effects specific fatty acids have on membrane structure and physiology are not well understood. However, the extraordinary selectivity of certain tissues for individual PUFAs strongly suggests a specific role for those fatty acids in cell function. For instance, brain and neural tissues are enriched in DHA, whereas adrenal glands have a high content of docosatetraenoic acid. Neither of these PUFAs can be produced de novo in humans and so their accumulation in cell membranes must be mediated by preferential absorption or specific desaturation reactions within the given tissue. PUFA specificity is also prevalent in subcellular organelles. Two examples include the inordinate enrichment of vaccenic acid in mitochondria and the conspicuous accumulation of arachidonic acid in the sn-2 position of nuclear membrane ether-linked phospholipids. Selective incorporation of PUFAs implies that fatty acids have some degree of functionality in the cell. In
addition, these functions must be monitored by the cell in order for there to be a regulatory adaptation. How a cell senses its acyl composition is not understood. To date, the best data on adaptive acyl regulation in mammalian cells involve the essential fatty acid deficiency response. Cells deficient in arachidonic acid or its precursor n-6 fatty acids convert de novo-produced oleic acid to n-9 eicosatrienoic (mead) acid by the action of the \( \Delta 5 \) and \( \Delta 6 \) desaturases. The conversion mechanism is identical to that of the production of C20 n-6 and n-3 fatty acids. The unusual desaturation of oleic acid is commonly believed to be the result of a cell’s attempt to replace arachidonic acid in the membrane with a similarly unsaturated species. This is an example of a cross-family compensation and provides further evidence for the lack of an association between n designation and function. The regulation of mead acid production is intriguing in that it is not found in animals fed sufficient dietary n-6 fatty acids. This may be the consequence of a differential activity of the \( \Delta 6 \) desaturase on linoleic and oleic acids.

Given that each fatty acid is unique in terms of its chemistry and potentially its function, one interesting question concerning PUFA metabolism is, can animals recognize n designation? If not, how big of a role do dietary PUFAs play in physiology? How can animals regulate the production of important acyl compounds in the face of changing dietary n-3/n-6 ratios, given that animals have a redundant system of desaturation for both n families? Changing n-3/n-6 PUFA ratios appears to have significant effects on much pathology in humans, including cardiovascular disease, immune function, and cancer. Because dietary acyl compositions vary tremendously throughout the world and over time, key questions must be answered concerning the regulatory factors and signals associated with the production of fatty acid and their metabolic products.

REFERENCES


I. EICOSANOIDS

There is a complex relationship among dietary fats, eicosanoids, and the immune system. Because long chain polyunsaturated fatty acids are the precursors for eicosanoids, these dietary components have the potential to modify levels of the products in the body. This is especially true if there is heavy reliance on a single fat in the diet. While this is the standard approach in nutrition studies in animals, it is also true that in a number of countries a single fat source provides as much as two-thirds of the human population’s total fat intake. In controlled feeding trials in humans, both the type and the amount of fat have been varied to permit an examination of the effects on eicosanoid production or changes in immune status.

Eicosanoids are only one of many possible mediators through which diet can influence the immune response. Eicosanoids are a large group of cyclized derivatives of the essential fatty acids that have potent biological activities and always contain 20 carbon atoms. These compounds usually have very short half-lives (measured in seconds) and are derived from the precursor fatty acids via a series of enzymatic steps (Fig. 1). In addition to the prostaglandins and leukotrienes, a variety of derivatives of 20-carbon fatty acids are produced. These include hydroxy and hydroperoxy fatty acids as well as hydroxylated or epoxydized derivatives. Cyclooxygenases add two oxygen molecules, lipoxygenase adds a single oxygen molecule, and cytochrome P450s add one atom of oxygen to the fatty acid.

The principal characteristic of the essential fatty acids (EFAs) is the presence of two or more cis double bonds in the families of ω3 or ω6 fatty acids, which must be derived from the diet. Fatty acids containing trans double bonds do not have EFA
activity and may increase the requirement for EFA. Endogenously synthesized unsaturated fatty acids are of the \( \omega 9 \) series and cannot be converted to the EFA precursors. EFA-deficient animals have an increase in 20:3 \( \omega 9 \) (derived from oleic acid) in tissue, and the triene/tetraene ratio in plasma lipids is often used clinically for diagnosis of this condition. This ratio essentially reflects the relative proportions of 20:3 \( \omega 9 \) to 20:4 \( \omega 6 \). Individuals with fat malabsorption (especially cystic fibrosis patients) or those on long-term parenteral nutrition are most likely to display signs of EFA deficiency. Diet-derived fatty acids are transported from the intestine to all tissues of the body, first via chylomicrons and then via other lipoproteins to all tissues in the body, usually becoming incorporated into the structurally important phospholipids of cell membranes.

Phospholipids of plasma membranes from mammals tend to be relatively enriched in long chain polyunsaturated fatty acids. Some of these serve as the precursors for synthesis of eicosanoids, principally arachidonic acid (20:4, \( \omega 6 \)), eicosapentaenoic acid (EPA, 20:5 \( \omega 3 \)) and dihomo-\( \gamma \)-linolenic acid (20:3, \( \omega 6 \)). Although \( \alpha \)-linolenic acid (18:3 \( \omega 3 \)) is the first fatty acid of the \( \omega 3 \) series, the biological effects of the 20-carbon fatty acids are usually stronger, presumably because of their direct steric competition with arachidonic acid as substrates for the enzymes of further metabolism. Most, if not all, terrestrial animals have limited ability to
synthesize EPA from linolenic acid except in tissues of the central nervous system. In contrast, fish are metabolically capable of these steps (they convert the 18:3 in plankton to 20:5), so they are a rich source. Another fatty acid concentrated in fish oil is docosahexaenoic acid (DHA, 22:6, ω3), which can potently inhibit cyclooxygenase. Thus, fish oil feeding has become an important paradigm in the study of the effects of dietary fat on eicosanoid metabolism and immune responses. It should be remembered that fish oils contain highly variable amounts of ω3 fatty acids, and many fish oils are high in saturated fatty acids [1].

Interest in eicosanoids, dietary fat, and the immune system is high because eicosanoid metabolism probably is one of the major mechanisms by which dietary fat modulates a variety of immune responses, although undoubtedly other mechanisms exist. Both the concentration of precursor fatty acid in the phospholipids of cell membranes and their rate of metabolism are controlling factors in the amount of eicosanoids released by cells. Competition between ω3 and ω6 fatty acids for lipoxygenases and cyclooxygenases results in production of eicosanoids of different types and in varying amounts. In the prostaglandin family there are two major series, designated E and F. The E series has a ketone and hydroxyl group added at C-9 and C-11, while the F series has hydroxyl groups at both positions, eliminating the double bonds from these positions (Table 1). In addition to the letter designation, there are subseries indicated by numbers. Those prostaglandins derived from 20:3 are designated E₁ or F₁; those from 20:4 are E₂ or F₂; and those derived from 20:5 are E₃ or F₃. Therefore, the subscripts 1, 2, and 3 refer to the number of double bonds remaining in a prostaglandin molecule.

The most common method for quantitating eicosanoids is through the radioimmunoassay (RIA). The RIA depends on an antibody that recognizes these molecules and does not cross-react with related compounds. Problems associated with the measurement of eicosanoids are the short half-life in tissues and body fluids as well as the cross-reactivity seen between some prostaglandins. For example, few methods can distinguish between prostaglandins E₁ and E₂, so they are often simply expressed as PGE. The short half-life of these compounds necessitates rapid isolation and chilling of samples, addition of a metabolic inhibitor, or all three. There is little doubt that many of the studies reporting levels of eicosanoids did not use conditions rigorous enough to ensure the accuracy of the analytical data. Moreover, since in-

Table 1  Relationship of Precursor Fatty Acids to Eicosanoid Products

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Prostaglandins</th>
<th>E series</th>
<th>F series</th>
<th>Thromboxanes</th>
<th>Leukotrienes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dihomo-γ-linolenic acid, all-cis-8,11,14 C20:3 (ω6)</td>
<td>E₁</td>
<td>F₁₉₅₆</td>
<td>A₁</td>
<td>A₁, C₁, D₁</td>
<td></td>
</tr>
<tr>
<td>Arachidonic acid, all-cis-5,8,11,14 C20:4 (ω6)</td>
<td>E₂</td>
<td>F₂₉₅₆</td>
<td>A₂</td>
<td>A₁, B₁, C₁, D₁, E₁</td>
<td></td>
</tr>
<tr>
<td>Eicosapentaenoic acid, all-cis-5,8,11,14,17 C20:5 (ω3)</td>
<td>E₃</td>
<td>F₃₉₅₆</td>
<td>A₃</td>
<td>A₃, B₃, C₃</td>
<td></td>
</tr>
</tbody>
</table>

Subscripts indicate the number of double bonds.

Can also be converted to prostacyclin (PGI₂).
stability is a characteristic of some eicosanoids, stable analogs may be measured in place of the compound of interest.

The leukotrienes are not cyclized and, as the name implies, contain three double bonds in the acyl chain. Arachidonic acid is metabolized by 5-lipoxygenase to leukotriene A₄. This eicosanoid can be metabolized by an epoxide hydrolase to leukotriene B₄ or via glutathione S-transferase to leukotriene C₄. The latter reaction results in the addition to the fatty acid of cysteine, glycine, and glutamic acid. Leukotriene C₄ can then be metabolized to D₄ by γ-glutamyltransferase, which removes the glutamic acid. Finally, D₄ can be metabolized to leukotriene E₄ by cysteinyl-glycine dipeptidase, which removes the glycine and leaves only the cysteine residue on the acyl chain. These compounds are critical mediators of a variety of inflammatory responses (e.g., anaphylaxis, increased vascular permeability, attraction of leukocytes and their activation).

Other eicosanoid products include thromboxanes, which are synthesized by platelets and cause platelet aggregation and vascular constriction. Finally, prostacyclin (PGI₂) is produced by blood vessels and inhibits platelet aggregation. These categories of eicosanoids do not appear to participate in regulation of immune responses; rather, they are altered in response to the dietary manipulation that affect the prostaglandins and leukotrienes.

Currently, there is considerable debate about the quantitative requirements for ω3 fatty acids; it is fairly well established that the (EFA) requirement of both rodents and humans is on the order of 0.5% of energy. The EFA requirement is satisfied primarily by intake of linoleic acid (there is relatively little arachidonic acid in commonly consumed foods) plus consumption of a smaller amount of linolenic acid or its fatty acid metabolites, which are concentrated in fish oils. Alterations in some immune functions can be seen with both low and high intake of EFA. This can be conceptualized as a curve with a flattened top (Fig. 2), in which the normal range of immune function is seen over a wide range of adequate EFA intake, whereas decreased immune responses are seen with either deficient or excessive intakes. The presumptive primary mechanisms are changes in the membrane microenvironment and/or precursors of the eicosanoids, although a number of other possible links have been suggested.

II. IMMUNE SYSTEM

A brief review of the components of the immune system will facilitate an understanding of the effects of dietary fat on the immune response. The immune system is the most dispersed network of cells in the body that are not in physical contact yet cooperate functionally. It is composed of both fixed and mobile cells that protect against invading organisms and the development of abnormal (malignant) cells. If, however, it overreacts to self, the result is autoimmunity; if it overreacts to nonself, there is an allergic response. Some of these immune cells have very short lives but others, which form the basis of immunological memory, exist for many years and perhaps for the lifetime of the host. Fixed cells of the reticuloendothelial system (RES) are found primarily in the liver, spleen, bone marrow, thymus, lymph nodes, lungs, and intestines. Mobile cells are found throughout the circulation in high numbers, which increase quickly in response to the presence of foreign antigens. It is
Figure 2 Immune responses as a function of dietary essential fatty acid. Note the broad and imprecise range of EFA adequacy for immune function; in particular, the upper limit of "normal" EFA varies for different immune functions.

estimated that 80% of mobile leukocytes are normally sequestered in organs of the RES, while the remainder float in the bloodstream.

Functionally, the components of the immune system are often divided into response elements. That is, humoral (or circulating) and cell-mediated immunity are the most commonly used concepts for describing immune functions, although there is communication between these facets of immune cells via a variety of soluble products called cytokines. In addition to the distinctions of humoral and cell-mediated immunity, a number of other components of the immune system play a role in protecting against pathogenic organisms. These include the polymorphonuclear leukocytes, which are divided into neutrophils, basophils, and eosinophils based on their staining characteristics in a smear of blood treated with Giemsa or Wright's stain. Neutrophils participate in acute inflammatory reactions against bacteria and other foreign bodies. Basophils and eosinophils are components of allergic reactions. Eosinophils are elevated also in response to parasitic infections.

There is a wide range of normal immune functions and a wide range of normal numbers of circulating immune cells. For example, the normal range of leukocytes in the blood of humans is 5000 to 10,000/mm$^3$. Within this range there are no discernible differences in susceptibility to infection. In fact, there is little increased risk of infection unless total white blood cells drop below 1000/mm$^3$.

Functional terms used to describe the immune system include natural (innate) and specific (acquired). Natural immunity is a result of molecules such as cytokines...
or complement, or a result of the function of natural killer (NK) cells or phagocytes, or barriers such as mucosal surfaces. The natural immune system is able to attack a substance to which it has had no previous exposure; subsequent exposure to that stimulus does not increase the response. On the other hand, the specific immune response recognizes a specific feature (antigen) of the foreign substance or cell and generates an immunologic memory, which amplifies responses during subsequent exposures. The specific immune system can recruit portions of the innate system to function simultaneously. Lymphocytes are the primary effectors of the specific immune response. These cells “direct” much of the immunologic activity by the production of cytokines of many types.

Cytokines are soluble protein molecules that function as messengers to other leukocytes. They are produced and secreted primarily by monocytes, and their main targets are lymphocytes and other monocytes. Some of the important cytokines include interleukins, interferons, and tumor necrosis factors (TNFs). There are at least 15 distinct forms of interleukins, 3 interferons, and 2 tumor necrosis factors.

A. Humoral Immunity

Humoral immunity is a result of the circulation in the plasma of antibodies and the activity of B lymphocytes. The B stands for bursa of Fabricius in birds, which is the immunological equivalent of the bone marrow in mammals. Humoral immunity characterizes the production of antibodies by the interaction of B lymphocytes with plasma cells. Circulating antibodies are the products of these cells, but they cannot be produced without interaction with T, or thymus-derived, cells. In addition to antibody production, the complement system is part of the immune response. Complement is a series of proteins in the plasma that participate in antigen-antibody reactions, displaying a wide range of actions ranging from aiding in phagocytosis by leukocytes to killing of tumor cells. Complement serves to stimulate various functions of the leukocytes.

B. Cell-Mediated Immunity

Cell-mediated immunity is based on the interactions of T lymphocytes and monocytic cells. T cells have been divided into many subclasses based primarily on cell surface receptors, which vary in relation to the functions these cells perform. The generally accepted classification of T cells is the CD (cluster designation or cluster of differentiation) scheme, which depends diagnostically on antibodies to distinguish these cellular sets of lymphocytes. Therefore, T cells can be classified as CD4+ and CD8+, but these cells are often referred to by their effector functions as helper and suppressor cells, respectively. In addition, there are NK cells, LAK (lymphokine-activated killer) cells, and lymphocytes responsible for delayed-type hypersensitivity.

The monocytes and their derived macrophages are cells that protect against certain bacteria and tumor cells. While monocytes circulate in the blood, once they have migrated into tissues they usually become fixed in place and are called macrophages. There are macrophages normally lining venules in the liver (Kupffer cells), spleen, and lymph nodes; these permanently fixed cells are termed histiocytes. The relative proportions of leukocytes in the blood differ across species. While the predominant circulating cells in rodents are lymphocytes, granulocytes are the most common in humans. This difference is relevant in comparing the experimental work...
done predominantly in rodent species with studies in humans. Macrophages are one of the most important sources of eicosanoids in the body, playing a major role in a variety of inflammatory and immunologic responses. Membranes of these cells are particularly enriched with arachidonic acid. Different inflammatory stimuli have the ability to induce secretion of eicosanoids of varied forms. Lymphocytes do not seem able to produce eicosanoids but can release arachidonic acid and certainly respond to the effects of these mediators. Many research studies describe use of mononuclear cells—these are mixed populations of lymphocytes and monocytes, usually obtained by differential centrifugation of blood or other biological fluids.

C. Eicosanoids and Immunity

Since eicosanoids are soluble mediators of the inflammatory response, their role is critical in many types of immune reaction. One of the most widely used drugs in the world, aspirin, is a cyclooxygenase inhibitor that reduces prostaglandin synthesis. The anti-inflammatory effect of aspirin and other nonsteroidal anti-inflammatory drugs (NSAIDs) is due to blocking of the metabolism of arachidonic acid by cyclooxygenase. Some animal studies suggest that high doses of other prostaglandin inhibitors, like indomethacin, may enhance some mononuclear cell dependent responses. There are limited reports that humans with deficient immune responses respond to indomethacin treatment whereas some normal individuals do not benefit from treatment with this drug and others show an increase in certain antibody responses [2]. The variation in response might be a function of which eicosanoid is involved in the altered immune responses. Burn injury leads to elevations of prostaglandin E and immune hyporesponsiveness. Administration of prostaglandin inhibitors lowers the PGE levels and partially corrects the immune response. NSAID administration appears to prevent colon cancer in both humans and experimental animals; in addition, prostaglandin synthesis is decreased by these drugs. It is unknown how these compounds inhibit the growth of colon cancer [3]. While there may be direct effects of these prostaglandin inhibitors on several aspects of colon carcinogenesis, including changes in cell metabolism, the cell cycle, and expression of tumor suppressor proteins, one cannot rule out alterations in immune response.

D. Noneicosanoid Mediators

Although it is clear that dietary fatty acids exert powerful effects on eicosanoids, which in turn profoundly modify some aspects of the immune response, alternate mechanisms by which dietary fat can alter immune responses have been proposed. These include changes in membrane microviscosity and dependent events, as well as the direct activation of protein kinase C (PKC) by arachidonic acid. The role of PKC is modulated by diacylglycerols (DAGs), whose affinity for PKC is altered by changes in the fatty acid moieties on the DAGs. Other potential mechanisms that have been implicated are activation of GTP-binding proteins by fatty acids and changes in phospholipase activity in the cell membrane. Finally, increased oxidation is suspected as one of the mediators of ω3 fatty acid effects on immune cells; an increase in dietary vitamin E that restored the in vitro responses of T cells from subjects fed fish oil was interpreted as supporting the possibility of such mediating activity [4]. Also, carotenoids exhibit both antioxidant and immunomodulatory roles that may be related. Feeding of carotenoids raises the number of circulating lym-
phocytes, increases proliferation of cytotoxic T cells, and enhances rejection of skin grafts in mice [5]. Similar data from humans were also seen: a low-carotenoid diet led to decreased delayed-type skin hypersensitivity reactions (mediated by T cells), and supplementation with β-carotene restored the skin reactions to normal [6].

Although a variety of cell-based mechanisms that do not involve eicosanoids have been proposed for modulating the immune response, perhaps some of the strongest evidence comes from studies of general nutritional status. Both overnutrition and energy restriction have profound effects on the immune system’s regulation. Overnutrition that results in obesity depresses the immune response, while energy restriction that avoids nutrient deficiencies enhances the immune response, particularly in older animals [7].

III. DIETARY LIPIDS

A. Cholesterol

Dietary cholesterol has been studied extensively for its effects on immune functions, primarily in animals and in vitro. A fair number of studies in humans have been conducted, however. The rationale for this work is that cholesterol and fatty acids in the cell membrane are the primary determinants of membrane microviscosity, which in turn controls a number of events at the cell surface, including enzymatic activities such as those of phospholipase and cyclooxygenase. There is considerable variation in serum cholesterol concentrations among humans, but two factors have contributed to the relatively small number of studies showing a positive correlation between dietary cholesterol and changes in the immune response. First, dietary cholesterol is relatively weak in its ability to increase serum cholesterol. Second, the populations studied have not exhibited a broad enough range of serum cholesterol concentrations to permit observation of large effects on immunological functions in vivo (although some studies have noted differential responses in vitro). Therefore, experimental manipulations in animals or in vitro that entailed extreme differences in exposure to cholesterol have resulted in significant effects on immunologic responses.

It is impossible to singularly characterize the effect of cholesterol on immunologic responses. This is because cholesterol is delivered to cells in lipoproteins, complex aggregates of cholesterol, triglyceride, phospholipid, and one or more apoproteins to a variety of immunologic cells. It has been shown that isolated apolipoprotein E (apoE) inhibits lymphocyte proliferation, but its effectiveness depends on interaction with cholesterol and phospholipid. Recent interest in the role of oxidized lipoproteins in the development of atherosclerosis suggests that some of the immunologic effects attributed to cholesterol may have been due to oxidized cholesterol molecules. In fact, there is considerable evidence that oxidized low density lipoprotein (LDL) or cholesterol reduces immune responses by lymphocytes.

The change in immunologic response elicited by elevated cholesterol concentrations depends both on the concentration of cholesterol to which cells are exposed and the type of immunologic response being measured. There is no single direction. Most studies in this area have looked at lymphocyte functions, and it can be concluded that results are primarily dependent on lipoprotein concentration [8].

Studies done in hypercholesterolemic rabbits and monkeys suggest that both B- and T-cell-dependent functions are elevated. However, hypercholesterolemic
guinea pigs showed increased immune function with a doubling of baseline serum cholesterol but a significant reduction when concentrations of cholesterol were elevated fourfold. This suggests that there may be species specificity as well as specific ranges of serum cholesterol levels that modulate immune responses. Most human studies of lymphocyte function, in vivo or in vitro, have shown decreased functions in the presence of high total or LDL cholesterol [9]. However, the total number of circulating T-lymphocyte subsets CD3⁺, CD4⁺, and CD8⁺ in hypercholesterolemic children correlates with LDL cholesterol concentrations over a threefold range [10]. Since T lymphocytes are found in significant numbers in atherosclerotic plaques, it is presumed that the elevated LDL levels are associated with both development of arterial lesions and changes seen with chronic inflammation.

Monocyte and macrophage functions are reduced in most studies that have examined the effect of excess cholesterol. Similarly, functions of the polymorphonuclear leukocyte have also been reduced in response to a surplus of cholesterol. The functions are, to a great extent, dependent on plasma membrane microviscosity; excess cholesterol incorporation will stiffen the membranes, thereby decreasing the ability of these cells to engulf microbes. A variety of other immune functions have been examined under the influence of different concentrations of cholesterol, and most studies show generally reduced immune cell responses; however, there is considerable disagreement in the research literature on the overall effect of cholesterol on immune reactions. Reports generally agree that there is reduced resistance to bacterial or viral infections, but some specific components of the immune system are suppressed while others are enhanced in the presence of hypercholesterolemia.

With the view that atherosclerosis shares many traits of a chronic inflammatory condition gaining widespread acceptance only in the late 1980s, the role of cholesterol in the human immune response as it relates to the development of arterial lesions has been of considerable interest. Since a large percentage of the fat-filled foam cells in atheroma are derived from macrophages, it is logical to assume that cytokines and classical inflammatory repair mechanisms are at work as part of the atherogenic process.

B. Fatty Acids

Just as cholesterol plays a major role in determining the physical state of the cell membrane, so do fatty acids. The more polyunsaturated a fatty acid molecule is, the more fluidity it imparts to the cell membrane. Also, the more polyunsaturated fat in the diet, the lower the immune response, within certain limits. That is, the immune response does not go down to zero in the presence of very high intake of polyunsaturated fat; it is, however, lower relative to that seen with ingestion of saturated fats. In both humans and animals, cell membrane phospholipids reflect dietary intake within limits of the membrane to accommodate a certain range of fatty acids. Dietary fat primarily alters storage of fatty acids in adipose tissue, but smaller changes in the membranes of many other cell types, including those of the RES, have been reported. A variety of experimental studies have demonstrated that both the amount and type of fatty acids consumed have roles to play in altering immune responses. Although it is easier to discuss specific fatty acids in the diet, it should be remembered that there are very few free fatty acids consumed and most are in triacylglycerol molecules, which are usually a mixture of two or three different fatty acids attached.
to a glycerol molecule. The majority of dietary fats usually supply four to six different fatty acids.

There is considerable evidence that both ω6 and ω3 polyunsaturated fatty acids (PUFAs) can modulate a variety of immunological activities. There was considerable interest in the clinical use of ω6-enriched vegetable oils during the 1970s for patients who had received kidney transplants, as well as those with multiple sclerosis and autoimmune disorders. Development of better immunosuppressive agents and other drugs for these conditions led investigators away from this area of research. However, there remains considerable interest in the effects of dietary fats on immune response, particularly in relation to development of cancer but also for cardiovascular disease and general health. It is fair to state that beyond the generally accepted immunosuppressive effects of polyunsaturated fats, there is considerable disagreement on whether the specific fatty acid, ratios between different fatty acids, or the quantity of dietary fat has the greatest impact on immunological responses. In general, when the long chain ω3 fatty acids increase in a cell membrane, there is a concomitant decrease in arachidonic acid concentration.

Sources of ω3 fatty acids used most commonly in studies of immune function include fish oils, evening primrose oil, and flaxseed oil. Plant-derived oils generally supply 18:3, while the fish oils have the longer chain fatty acids EPA and DHA. There is substantial debate concerning the optimal range of the ω6/ω3 ratio in the diet with many researchers favoring a range of 1.0–7.0 based on studies in rats. The 10th edition of the U.S. Recommended Dietary Allowances does not make specific recommendations for any essential fatty acids in humans but indicates the desirability of establishing these in the near future.

One study of the immune response in rats fed blends of different proportions of sunflower oil (rich in linoleic acid) and flaxseed oil (rich in linolenic acid) found that the higher the ω3/ω6 ratio of fatty acids in the plasma, the greater the reduction of lymphocyte-dependent immune responses, including T-cell blastogenesis and natural killer cell activity [11]. These authors concluded that α-linolenic acid was as potent as fish oil for suppressing immune responses.

While most animal studies simply use feeding of a single source of fat, or a blend, throughout an experiment, at least one study has tried to mimic the normal human intake pattern of ω3 PUFA [12]. Mice were fed a diet containing safflower oil and switched to a sardine/olive oil mixture for periods of 1–7 days per week. Peritoneal macrophages had phospholipid compositions that reflected the diet. Synthesis of leukotrienes E₄ and C₄ decreased, while E₅ and C₅ levels increased with more frequent consumption of fish oil. Prostaglandin F₁₉ also decreased with increasing consumption of fish oil. The summary finding of this study—that fish oil must be consumed at least twice a week to produce significant changes from the control diet—touches on the current debate about how frequently a person has to consume fish to derive a health benefit. As with most questions on diet and risk of chronic diseases, there is no definitive answer, but some studies have suggested maximal benefit with consumption of at least two servings of fatty fish weekly.

A fundamental study on this topic looked at the effects of feeding a low-fat (26% of energy) diet that contained a high amount of fish in 22 subjects for 24 weeks or a low-fat diet without fish [13]. Fish intake ranged from 4 to 6 ounces daily. Therefore, this study used both type and amount of dietary fat as variables. Responses on these diets were compared with those of subjects who followed a diet with 35%
of energy from fat. Feeding the low-fat, low-fish diet (which was enriched in plant PUFAs) increased the response of mononuclear leukocytes to the T-cell mitogen concanavalin A (con A), interleukin 1β levels, and tumor necrosis factor. No effects were seen on PGE₂ production, interleukin 6 levels, or delayed-type skin hypersensitivity. In contrast, the low-fat, fish-enriched diet resulted in a significant decrease in CD4⁺ and a concurrent decrease in CD8⁺ cells. There were significant reductions in the lymphocyte mitogenic response to con A, delayed-type hypersensitivity, interleukin 1β, interleukin 6, and tumor necrosis factor production by leukocytes. The practical implications of these immune alterations are not clear cut. It was conjectured that these decreases in immune responses would be favorable for atherosclerosis and inflammatory diseases but harmful for host defense against microorganisms. The issue of immune surveillance against cancer cells was not discussed but is a valid concern.

In another important study in this area, Purasiri et al. [14] found that ω3 fat supplement of 4.8 g daily decreased the blood concentrations of a number of cytokines significantly (by 60–80%) in individuals with colon cancer; these included several forms of interleukins, tumor necrosis factor α, and interferon-γ. Cytokines returned to baseline levels 3 months after the cessation of the supplements. This study clearly demonstrated a rapid and profound decline in a number of cytokines in response to an easily consumed amount of ω3 fatty acids. The implications of this study are that regular consumption of ω3 fatty acids might reduce the ability of the immune system to respond to infections or tumors.

Some studies have measured prostaglandin metabolites in urine as an index of whole-body metabolism. Supplementation of ω3 fatty acids to a low-fat diet reduced thromboxane A₂ production, while a low-fat diet, with or without fish oil, resulted in reduced prostacyclin production in healthy men. Accompanying these dietary manipulations, however, were increases in thromboxane A₁ and prostacyclin A₁ [15].

There have been many studies in which fish oil supplements were administered to people with presumptive autoimmune inflammatory conditions such as rheumatoid arthritis. Most studies find that fish oil supplements are effective in reducing symptoms of these conditions. However, there does not seem to be any benefit of ω3 fatty acid supplements over conventional NSAID therapy.

C. Total Dietary Fat

The early work on total dietary fat in the 1960s showed that animals fed high-fat diets were more susceptible to a wide range of spontaneous infections. These observations led to the experimental examination of specific parts of the immune response in animals fed different levels of dietary fat.

Differences in immune responses as a function of changes in total dietary fat have been studied in many animal models and in a few human trials. In general, the more fat in the diet, the lower the immune response. Many animal studies have shown high-fat diets are immunosuppressive in comparison to low-fat diets. The basal level in most animal studies have been about 10% of energy from fat, and the high levels have ranged from 40 to 60% of energy from fat, usually of a single source. Most animal studies have fed a polyunsaturated vegetable oil, so it is difficult to distinguish between the effects of total fat and the effect of increased linoleic acid or other individual fatty acids.
A limited number of published human studies have examined the influence of amount of dietary fat on immune responses. In one experiment, seven healthy women lived in a metabolic suite and were fed a diet that provided 41% of energy from fat and 5% from PUFAs [16]. The subjects were divided into two groups that consumed either 26% of energy from fat with 3.2% from PUFAs or 31% of energy from fat with 9.1% PUFAs; the study was conducted in a crossover design. Both low-fat diets resulted in significant increases in serum complement fractions C3 and C4, as well as mitogenic responses of peripheral lymphocytes, phytohemagglutinin, con A, protein A, and pokeweed mitogen. The results are similar to those of Meydani et al. [13], cited above, namely, that lower fat diets increased blastogenic response of blood lymphocytes to con A. In another study, healthy women were fed either 3% or 8.3% of energy as linoleic acid [17]; increasing dietary 18:2 led to higher prostaglandin levels in urine but lower thromboxane B2 and no change in prostaglandin F1α.

REFERENCES


I. INTRODUCTION

Many factors are associated with increased risk for coronary heart disease (CHD), a major cause of morbidity and mortality in prosperous Western countries. Some of these factors, such as increasing age or a family history of premature CHD, are not amenable to preventive intervention, but other factors are. Three of these preventable factors—the distribution of plasma cholesterol over the low density and high density lipoproteins (LDLs and HDLs), the oxidizability of LDLs, and hemostasis—can be modified by changing the sources and amount of fats and oils in the diet. The purpose of this chapter is to review some of the most recent and important findings on the effects of dietary fatty acids on these three risk factors.

II. DIETARY FATS

Although fat and oils are complex mixture of fatty acids, each fat or oil has its characteristic fatty acid composition. Dairy fat, for example, is relatively rich in fatty acids with 14 or less carbon atoms, whereas olive oil has a high oleic acid content (Table 1). Sunflower oil, on the other hand, is rich in linoleic acid, although certain varieties exist that contain large amounts of oleic acid. In normal, regular diets, palmitic and stearic acids are the most prevailing saturated fatty acids, while oleic and linoleic acids are, respectively, the most widespread dietary monounsaturated
Table 1  Major Fatty Acids in Some Edible Fats and Oils

<table>
<thead>
<tr>
<th>Formula</th>
<th>Fatty acid</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saturated fatty acids</td>
<td>Medium chain fatty acids</td>
<td>Dairy fat, coconut oil, palm kernel oil</td>
</tr>
<tr>
<td>MCFA</td>
<td>C12:0 Lauric acid</td>
<td>Dairy fat, coconut oil, palm kernel oil</td>
</tr>
<tr>
<td>C14:0 Myristic acid</td>
<td>Dairy fat, coconut oil, palm kernel oil</td>
<td></td>
</tr>
<tr>
<td>C16:0 Palmitic acid</td>
<td>Palm oil, meat</td>
<td></td>
</tr>
<tr>
<td>C18:0 Stearic acid</td>
<td>Meat, cocoa butter</td>
<td></td>
</tr>
<tr>
<td>Monounsaturated fatty acids</td>
<td>C18:1,n-9 Oleic acid</td>
<td>Olive oil, rapeseed oil, high oleic acid sunflower oil</td>
</tr>
<tr>
<td>Polyunsaturated fatty acids</td>
<td>C18:2,n-6 Linoleic acid</td>
<td>Sunflower oil, corn oil, soy bean oil, corn oil</td>
</tr>
<tr>
<td>C18:3,n-3 α-Linolenic acid</td>
<td>Rapseed oil, soy bean oil</td>
<td></td>
</tr>
<tr>
<td>C20:5,n-3 Tammnodonic acid</td>
<td>Fatty fish, fish oil capsules</td>
<td></td>
</tr>
<tr>
<td>C22:5,n-3 Cervonic acid</td>
<td>Fatty fish, fish oil capsules</td>
<td></td>
</tr>
</tbody>
</table>

*The trivial name for tammnodonic acid is eicosapentaenoic acid (EPA) and for cervonic acid docosahexaenoic acid (DHA).

and polyunsaturated fatty acids. About 30–40% of total energy intake is provided by fat. For a person consuming 10 MJ (2400 kcal) per day, this corresponds to a fat intake of 80–107 g.

III. LIPOPROTEINS

The solubility of cholesterol in water is very low, approximately $5.2 \times 10^{-3}$ mmol/L. The actual cholesterol concentration in the watery plasma of healthy subjects, however, is about 3.9–5.2 mmol/L, and increases to more than 10 mmol/L in hypercholesterolemic people. This high degree of solubilization is achieved by the formation of lipoproteins.

Lipoproteins are globular, high molecular weight particles that are complex aggregates of lipid and protein molecules. A lipoprotein consists of a hydrophobic core, which mainly contains triacylglycerols and cholesterylesters, and a polar, hydrophilic coat composed of phospholipids, unesterified cholesterol, and specific apolipoproteins. In this way, the hydrophobic core is protected from the watery surrounding and transport of large amounts of cholesterol and triacylglycerols through the blood vessels is possible.

Lipoproteins are a heterogeneous group, which can be divided into five major classes: chylomicrons, very low density lipoproteins (VLDLs), intermediate density lipoproteins (IDLs), low density lipoproteins (LDLs), and high density lipoproteins.
(HDLs). Each class has its own characteristic lipid and apolipoprotein composition, size, and density, whereas each apolipoprotein has its own specific metabolic functions (Tables 2 and 3).

**IV. METABOLISM OF DIETARY FATTY ACIDS AND LIPOPROTEINS**

**A. Exogenous Pathway**

The metabolism of dietary fatty acids, dietary cholesterol, and lipoproteins is depicted in Figure 1. In the duodenum, dietary triacylglycerols are dissolved with the help of bile salts, as well as small quantities of fatty acids and monoglycerides. The enzyme pancreatic lipase then hydrolyzes the dietary triacylglycerols into mono- and diglycerides, free fatty acids, and glycerol. The so-formed emulsion of lipids, which also contain the dietary cholesterol, passes the mucous membrane of the intestinal cells. Within the cell, further hydrolysis of lipids takes place and new triacylglycerols are formed by reesterification of the free fatty acids with glycerol. The newly synthesized triacylglycerols and cholesteryl esters, derived from the dietary cholesterol, are incorporated into chylomicrons, which enter the lymph and subsequently the blood circulation in the subclavian vein. In the blood, the triacylglycerols from the chylomicron core are hydrolyzed by lipoprotein lipase (LPL), an enzyme adhered to the endothelial cells of the blood vessels and activated by apolipoprotein C-II. The free fatty acids pass the endothelial cells and enter adipocytes or muscle cells. In these cells, the fatty acids are respectively stored as triacylglycerols or oxidized. The core of an emptied chylomicron mainly consists of cholesteryl esters and is called a *chylosmicron remnant*. These remnant particles are removed from the circulation by the hepatic remnant receptor, which has a high affinity for apolipoprotein E (apoE) from the chylomicron surface.

**Table 2** Some Physical Characteristics and Mean Composition of Lipoprotein Fractions from Normotriglyceridemic Subjects

<table>
<thead>
<tr>
<th></th>
<th>Chylomicrons</th>
<th>VLDL</th>
<th>IDL</th>
<th>LDL</th>
<th>HDL</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Density (g/mL)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lower limit</td>
<td>—</td>
<td>0.96</td>
<td>1.006</td>
<td>1.029</td>
<td>1.063</td>
</tr>
<tr>
<td>Upper limit</td>
<td>0.96</td>
<td>1.006</td>
<td>1.019</td>
<td>1.063</td>
<td>1.21</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Core components</th>
<th></th>
<th>Surface components</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>% of total lipoprotein mass</strong></td>
<td>Triacylglycerols</td>
<td>87</td>
<td>Phospholipids</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>Cholesteryl esters</td>
<td>3</td>
<td>Free cholesterol</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Apolipoprotein</td>
<td>2</td>
<td></td>
<td>2</td>
</tr>
</tbody>
</table>

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Table 3  Some Major Apolipoprotein and Their Functions

<table>
<thead>
<tr>
<th>Apolipoprotein</th>
<th>Lipoprotein</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>ApoC-II</td>
<td>Chylomicrons, VLDLs</td>
<td>Activator of LPL</td>
</tr>
<tr>
<td>ApoE</td>
<td>Chylomicron remnants, VLDLs, IDLs</td>
<td>Ligand for remnant and LDL receptors</td>
</tr>
<tr>
<td>ApoB-100</td>
<td>IDLs, LDLs</td>
<td>Ligand for LDL receptor</td>
</tr>
<tr>
<td>ApoA-I</td>
<td>Chylomicron, HDLs</td>
<td>Cofactor for LCAT</td>
</tr>
</tbody>
</table>

B. Endogenous Pathway

The liver excretes cholesterol and triacylglycerols into the circulation by the formation of VLDL particles. These lipoproteins also interact with LPL, triacylglycerols become hydrolyzed, and VLDL remnants (also called IDLs) are formed. A part of the IDL is taken up by the liver, while the remaining part is converted in the circulation into LDL. LDLs are nearly devoid of triacylglycerols and carry about 60-
70% of the total amount of cholesterol in the plasma. Most of the LDL is now removed from the circulation through the hepatic LDL receptor pathway, which recognizes apolipoprotein B-100 (apoB-100). LDL uptake from the blood by the LDL receptor–mediated pathway is highly controlled and this pathway will be down-regulated if the amount of cholesterol in the cell becomes too high. A smaller part, however, is removed via the scavenger pathway. Uptake via this pathway is not saturable and is positively related with the LDL cholesterol concentration. Thus, the higher the blood LDL cholesterol concentration, the more LDL will be taken up via the scavenger pathway [1]. When too much LDL is taken up via the scavenger pathway from macrophages, cells loaded with cholesterol are formed—so-called foam cells—which are frequently found in atherosclerotic lesions.

Cholesterol can also be transported out of tissues by reverse cholesterol transport. This system is mediated by HDL, lecithin cholesteryl acyltransferase (LCAT) and cholesterol ester transfer protein (CETP). HDL binds free cholesterol from tissues, which is esterified by LCAT, a protein associated with apolipoprotein A-I (apoA-I). The formed cholesterylesters move to the core of the HDL particle and the HDL is converted to a larger particle. The acquired cholesterylesters can now be transferred with the assistance of CETP to apoB-100-containing lipoproteins in exchange for triacylglycerols. The apoB-100-containing lipoproteins are further metabolized, as has already been described. The large HDL particles may also be taken up in the liver by a putative HDL receptor or lose a part of its content by the action of hepatic lipase and then reenter the circulation again.

V. PLASMA LIPOPROTEINS AND CORONARY HEART DISEASE

LDL and HDL have different effects on the risk for CHD. High concentrations of LDL are atherogenic, whereas high levels of HDL are negatively associated with the risk for CHD. As LDL carries most of the plasma cholesterol, the total plasma cholesterol is also a good index for the risk of CHD. It should be realized, however, that some people have high total cholesterol concentrations, due to high HDL levels. Therefore, the total cholesterol to HDL cholesterol might be the most efficient predictor for the risk for CHD [2]. Also, high levels of triacylglycerols, which are in the fasting condition mainly found in the VLDLs, are positively related to the risk for CHD [3].

VI. DIETARY FATS AND PLASMA LIPOPROTEINS

A. Earlier Studies

In the 1950s Keys and coworkers started a series of well-controlled experiments to examine the effects of dietary fatty acids on plasma total cholesterol concentrations [4,5]. Groups of physically healthy men were fed diets that differed widely in the amount of fat and in dietary fatty acid composition. During the studies, individual allowances were adjusted weekly to keep body weight stable so that changes in plasma total cholesterol concentrations could be attributed solely to dietary changes. At the end of the studies, an empirical formula was derived that could be used to predict for a group of subjects changes in plasma cholesterol concentrations from changes in dietary fatty acid composition:
\[ \Delta \text{ plasma total cholesterol (mmol/L)} = 0.03 \times (2 \times \Delta \text{Sat'} - \Delta \text{Poly}) \]

or

\[ \Delta \text{ plasma total cholesterol (mg/dL)} = 1.2 \times (2 \times \Delta \text{Sat'} - \Delta \text{Poly}) \]

where Sat’ are the percentages of energy provided by saturated fatty acids with 12, 14, or 16 carbon atoms (lauric, myristic, and palmitic acid, respectively), whereas Poly refers to the amount of polyunsaturated fatty acids in the diet.

How should this formula be interpreted? First, it should be realized that effects are expressed relative to those of carbohydrates. A hypercholesterolemic fatty acid is therefore defined as a fatty acid that causes an increase in the plasma cholesterol level when substituted in the diet for an isocaloric amount of carbohydrates. Thus, when 10% of energy from Sat’ is replaced by carbohydrates, \( \Delta \text{Sat’} \) equals \(-10\), and the expected decrease in plasma total cholesterol concentrations is

\[ 0.03 \times 2 \times -10 = -0.60 \text{ mmol/L} \left[ -24 \text{ mg/dL} \right] \]

If this amount of carbohydrates is then replaced by linoleic acid, a further decrease of

\[ -0.03 \times 10 = -0.30 \text{ mmol/L} \left[ -12 \text{ mg/dL} \right] \]

is expected. Direct replacement of Sat’ by Poly will yield the sum of these two effects, a fall of 0.90 mmol/L [36 mg/dL]. Further, this formula suggests that—because they are not part of the equation—the effects on plasma total cholesterol concentrations of saturated fatty acids with fewer than 12 carbon atoms, of stearic acid, and of monounsaturated fatty acids are similar to those of carbohydrates. Finally, it can be seen that the cholesterol raising effect of Sat’ is about twice the cholesterol lowering effect of Poly.

Similar types of studies were carried out in the 1960s by Hegsted and colleagues [6]. Results were essentially similar, but it was also concluded that myristic acid was more cholesterolemic than palmitic and lauric acids.

These and other studies have lead to recommendations that the most effective diet for lowering plasma total cholesterol concentration should contain a low proportion of the cholesterol-raising saturated fatty acids and a high proportion of linoleic acid. Also, a reduction in cholesterol intake was advocated as dietary cholesterol increases plasma total cholesterol concentrations [6,7].

However, these earlier well-controlled studies were not specifically designed to examine the effects of specific dietary fatty acids on plasma cholesterol concentrations and over the various lipoproteins. Therefore, new studies were initiated that compared side-by-side effects of specific fatty acids on the plasma lipoprotein profile.

**B. Recent Studies**

1. **Saturated Fatty Acids**

To discuss the effects of saturated fatty acids on plasma lipid and lipoproteins, the saturated fatty acids, in agreement with the results of Keys and colleagues [5], fall into three classes: medium chain fatty acids; fatty acids with 12, 14, or 16 carbon atoms; and stearic acid.

a. **Medium Chain Fatty Acids.** Saturated fatty acids with fewer than 12 carbon atoms are called short and medium chain saturated fatty acids (MCFAs) and are found in relatively large amounts in coconut fat, palm kernel oil, and butterfat, but also in certain structured lipids, parenteral nutrition preparations, and sport drinks.
McGandy and coworkers [8] have carefully compared the effects of MCFAs, mainly capric acid, on plasma total and LDL cholesterol levels. Eighteen physically healthy men were fed several diets, each for 4 weeks. Diets contained several low-fat food items to which the experimental fats were added. It was shown that modest amounts of MCFAs in the diet have comparable effects on the plasma total and LDL cholesterol and on triacylglycerols concentrations as have carbohydrates. However, large amounts of MCFAs increased triacylglycerol concentrations. Recent results of Cater et al. [9] suggested that a mixture of MCFAs slightly decreases LDL cholesterol concentrations relative to palmitic acid but increases LDL cholesterol relative to oleic acid (Fig. 2).

b. Lauric and Myristic Acids. Palm kernel oil, coconut oil, and dairy fat are rich in lauric acid but also contain relatively high amounts of myristic acid. Therefore, it is hardly possible to study the specific effects of these two saturated fatty acids on plasma lipoproteins with natural fats. For example, a diet enriched in palm kernel oil will contain high amounts of both lauric and myristic acids, and it will subsequently be impossible to ascribe the effects on the plasma lipoprotein profile to lauric or myristic acids. To circumvent this problem, most studies have used synthetic fats to examine the cholesterolemic effects of these two saturated fatty acids. In this way, a fat with any desired fatty acid composition can be made. For example, when one interesterifies trilaureate with a high oleic acid sunflower oil, the result will be a high lauric acid fat without any myristic acid.

Denke and Grundy [10] compared the effects on plasma lipoproteins of lauric acid with those of palmitic acid and oleic acid. Three different liquid formula diets, which differed only in the type of fat used, were fed for 3 weeks in random order to 14 men. It was concluded that lauric acid raised plasma total and LDL concentrations as compared with oleic acid, but to a lesser extent than the palmitic acid.

![Figure 2](image.png)

**Figure 2** Effects of medium chain triacylglycerols and palmitic acid on plasma total, LDL cholesterol, HDL cholesterol, and triacylglycerol concentrations relative to those of cis-monounsaturated fatty acids (oleic acid). Nine men received three mixed natural diets, each for 3 weeks, in random order. The composition of the diets was identical, except for 43% of daily energy intake, which was provided as medium chain triacylglycerols (C8:0 and C10:0), palmitic acid (C16:0), or oleic acid (cis-C18:1) [9].
diet did. No effects were observed on plasma HDL cholesterol or triacylglycerol concentrations. Using mixed natural diets, however, Temme et al. have found that lauric acid raised total cholesterol concentrations more than palmitic acid, which was partly due to an increase in HDL cholesterol [11] (Fig. 3).

The effects of myristic acid have been examined in several studies with mixed solid foods. Zock et al. [12] concluded that myristic acid increased plasma total cholesterol concentrations relative to palmitic and oleic acids, which was due to an increase in both LDL and HDL cholesterol concentrations. However, triacylglycerol concentrations were not affected (Fig. 4). Tholstrup et al. [13] also found that myristic acid increased HDL cholesterol concentrations compared with palmitic acid. However, no differences were observed in plasma LDL cholesterol concentrations.

c. Palmitic Acid. Replacing palmitic acid in the diet for carbohydrates lowers LDL and HDL cholesterol concentrations. However, if the intake of palmitic acid is decreased at the expense of oleic acid, LDL cholesterol concentrations decrease, whereas HDL cholesterol is not affected [10–12] (Figs. 2–5). Recently, it has been postulated that in normolipidemic, nonobese subjects palmitic acid may not always be a cholesterol-raising saturated fatty acid [14,15], provided that linoleic acid intake contributes at least 6–7% of daily energy intake and daily dietary cholesterol intake is less than 300 mg. Of course, these findings will be of great practical significance if it proves to be correct that under certain conditions palmitic acid can be exchanged for oleic acid without affecting LDL cholesterol levels. However, these studies [14,15] await confirmation.

d. Stearic Acid. Keys and coworkers already demonstrated that stearic acid did not increase plasma total cholesterol concentrations [5,16]. However, the effects of stearic acid on the distribution of cholesterol over the various lipoproteins were
Figure 4  Effects of myristic and palmitic acids on plasma total, LDL cholesterol, HDL cholesterol, and triacylglycerol concentrations relative to those of cis-monounsaturated fatty acids (oleic acid). Thirty-six women and 23 men received three mixed natural diets, each for 3 weeks, in random order. The composition of the diets was identical, except for 10% of daily energy intake, which was provided as myristic acid (C14:0), palmitic acid (C16:0), or oleic acid (cis-C18:1) [12].

Figure 5  Effects of palmitic and stearic acids on plasma total, LDL cholesterol, HDL cholesterol, and triacylglycerol concentrations relative to those of cis-monounsaturated fatty acids (oleic acid). Eleven men received three liquid-formula diets, each for 3 weeks, in random order. The composition of the diets was identical, except for 16% of daily energy intake, which was provided as palmitic acid (C16:0), stearic acid (C18:0), or oleic acid (cis-C18:1) [17].
studied in detail for the first time by Bonanome and Grundy [17]. From that study it was concluded that stearic acid exerted similar effects on the plasma lipoprotein profile as oleic acid, a monounsaturated fatty acid. However, palmitic acid was clearly hypercholesterolemic relative to stearic acid (Fig. 5).

2. Monounsaturated Fatty Acids

The most abundant monounsaturated fatty acid in the human diet, oleic acid, has 18 carbon atoms and one double bond. Although olive oil probably is the most well-known source of oleic acid, animal fats are in many countries a major contributor to total oleic acid, but also to palmitic and stearic acid, intake.

a. Oleic Acid. Effects of monounsaturated fatty acids on plasma total cholesterol levels are often described as neutral. This term is often misinterpreted. It does not mean that the plasma total cholesterol level does not change when monounsaturates are added to the diet. Neutral indicates that monounsaturated fatty acids have the same effect on plasma total cholesterol as compared with an isocaloric amount of carbohydrates. Although this may be correct for plasma total cholesterol levels, oleic acid and carbohydrates do not have similar effects on the distribution of cholesterol of the various lipoproteins [18] (Fig. 6). This was also shown in a study with young, healthy volunteers [19]. Increasing the intake of oleic acid at the expense of carbohydrates increased plasma HDL cholesterol concentrations and decreased those of triacylglycerols. The increase in HDL cholesterol was compensated for by a decrease in VLDL cholesterol. Effects of carbohydrates and oleic acid on plasma total and LDL cholesterol concentrations were comparable.

![Figure 6](image-url) Effects of carbohydrates and palmitic acid on plasma total, LDL cholesterol, HDL cholesterol, and triacylglycerol concentrations relative to those of cis-monounsaturated fatty acids (oleic acid). Seven men received three liquid formula diets, each for 3 weeks, in random order. The composition of the diets was identical, except for 19% of daily energy intake, which was provided as carbohydrates, palmitic acid (C16:0), or oleic acid (cis-C18:1) [18].
3. Polyunsaturated Fatty Acids

Polyunsaturated fatty acids in the diet belong to either the n-6 or the n-3 family. About 90% of all polyunsaturated fatty acid in the diet is linoleic acid, which is found in vegetable oils like sunflower oil, corn oil, and soybean oil. The mean daily intake of fatty acids from the n-3 family is only 1–3 g. These polyunsaturated fatty acids are either from vegetable or animal origin. α-Linolenic acid is found in rapeseed and soybean oils, while the very long chain fatty acids timnodonic or eicosapentaenoic acid (EPA) and cervonic or docosahexaenoic acid (DHA) are only present in fish oils.

a. n-6 Polyunsaturated Fatty Acids. Earlier studies found that linoleic acid was hypocholesterolemic as compared with carbohydrates and monounsaturated fatty acids [5,6]. The study of Mattson and Grundy [20], however, suggested that, as compared with monounsaturates, part of the cholesterol-lowering effect of linoleic acid was due to a decrease in HDL cholesterol (Fig. 7). However, linoleic acid intake in that study was unrealistically high (28% of energy intake), which may have influenced the results. Studies at lower intakes found similar effects of linoleic and oleic acids on HDL cholesterol, but also on LDL cholesterol [21–23]. Thus, these more recent studies suggested that replacement of saturated fatty acids in the diet by monounsaturated fatty acids causes the same favorable change in plasma lipoprotein cholesterol levels as replacement by polyunsaturated fatty acids. However, it should be noted that other recent studies found a small favorable effect linoleic acid on plasma LDL cholesterol as compared with oleic acid [24].

b. n-3 Polyunsaturated Fatty Acids. The effects of α-linoleic acid on the plasma lipoprotein profile are similar to those of linoleic acid [25]. The highly unsaturated

![Figure 7](image)

**Figure 7** Effects of palmitic acid and linoleic acids on plasma total, LDL cholesterol, HDL cholesterol, and triacylglycerol concentrations relative to those of cis-monounsaturated fatty acids (oleic acid). Twelve men received three liquid formula diets, each for 4 weeks, in random order. The composition of the diets was identical, except for 14% of daily energy intake, which was provided as palmitic acid (C16:0), linoleic acid (C18:2), or oleic acid (cis-C18:1) [20].
fatty acids from fish oils, however, have different effects. In normocholesterolemic subjects, these fatty acids do not change plasma LDL or HDL cholesterol concentrations, but do lower plasma triacylglycerols and the concentration of cholesterol in VLDL [26]. In hyperlipidemic subjects, and in particular in patients with elevated triglyceride concentrations, fish oils also lower plasma triacylglycerols, but raise LDL and HDL cholesterol concentrations [27].

4. trans Fatty Acids

Most unsaturated fatty acids found in nature have the cis configuration. This means that the two carbon side chains attached to the double bond point into the opposite (cis) direction. However, in some fatty acids the carbon side chain point into the same (trans) direction. In this way, two compounds are formed that have exactly the same number and type of atoms, but have different chemical, physical, and physiologic characteristics.

trans Fatty acids are formed when vegetable oils are hardened by hydrogenation. These hydrogenated fats are used for the production of certain types of margarines, frying fats, and foods prepared with these fats. Most trans fatty acids in the diet have 18 carbon atoms and one double bond (trans-C18:1). However, trans fatty acids are not only found in hydrogenated oils but also in milk fat and body fat from ruminants, formed from dietary polyunsaturated fatty acids by the action of bacteria in the rumen of these animals. The trans polyunsaturated fatty acids in the diet mainly originate from hydrogenated fish oils.

a. trans Fatty Acids. LDL cholesterol concentrations increase when cis-monounsaturated fatty acids in the diet are replaced by trans-monounsaturated fatty acids [28]. In most studies, a decrease in HDL cholesterol was also observed. Although the LDL cholesterol raising effect of trans-monounsaturated fatty acids is less than the effect of a mixture of saturated fatty acids, trans-monounsaturated fatty acids also lowered HDL cholesterol relative to a mixture of saturated fatty acids. Therefore, it was concluded that both types of fatty acids have an unfavorable effect on the plasma lipoprotein profile [29] (Fig. 8).

C. Conclusion

Dietary fatty acid composition affects the distribution of cholesterol over LDL and HDL. As compared with an isoenergetic amount of carbohydrates, myristic acid (C14:0) is the most potent plasma total and LDL cholesterol–raising saturated fatty acid. From the other saturated fatty acids in the diet, lauric acid (C12:0) and, to a lesser degree, palmitic acid (C16:0), also have a hypercholesterolemic effect, whereas stearic acid (C18:0) seems to be neutral. Linoleic (cis,cis-C18:2) and probably also oleic acid (cis-C18:1) have a small LDL cholesterol–lowering effect. trans Fatty acids have a strong plasma total and LDL cholesterol–increasing effect. Effects on LDL cholesterol levels are positively related to those on HDL cholesterol concentrations. An exception are trans fatty acids that do not have an effect on HDL as compared with carbohydrates. Thus, a reduction in the intake of the cholesterol-raising saturated fatty acids and trans fatty acids is more important for optimizing the plasma lipoprotein profile than a reduction in total fat intake per se.
Figure 8  Effects of a mixture of saturated fatty acids and trans isomers of oleic acid on plasma LDL cholesterol, HDL cholesterol, and triacylglycerol concentrations relative to those of cis-monounsaturated fatty acids (oleic acid). Twenty-five men and 34 women received three mixed natural diets, each for 3 weeks, in random order. The composition of the diets was identical, except for 10% of daily energy intake, which was provided as saturated fatty acids (mainly lauric and palmitic acids (C12:0 and C16:0, respectively), trans isomers of oleic acid (trans-C18:1), or oleic acid (cis-C18:1) [28].

VII. OXIDIZABILITY OF LDL

As has already been mentioned, elevated plasma LDL cholesterol concentrations are associated with increased risk for CHD. However, the atherogenicity of the LDL particle increases after oxidative modification of its polyunsaturated fatty acids.

A. LDL Oxidation

Oxidation of LDL is a free radical–driven process that may initiate a cascade of reactions (Fig. 9). For the in vivo situation it is not clear where the initiating radical species is derived from, but several suggestions, based on in vitro experiments, have been made.

Some experiments have suggested that cellular production of superoxide anions ($O_2^-$) or hydroxyl radicals (OH$^-$), which are intermediates in several metabolic processes from the mitochondrial respiratory chain or the cytochrome P450 system, initiate the lipid peroxidation reaction. Other experiments have proposed that lipid peroxidation is initiated by lipoxygenase activity, as this enzyme forms radicals as intermediate products in the formation of eicosanoids. These hypotheses are not necessarily contradictory because lipoxygenase activity might be particularly important in endothelial cells and peroxide initiation by superoxide anions and hydroxyl radicals in smooth muscle cells [30].

In vitro oxidation of LDL results in alterations in both the lipid and the protein component of LDL. The amount of unsaturated cholesteryester content decreases, especially cholesteryl arachidonate and cholesteryl linoleate. In addition, phosphatidylcholine—the main phospholipid in LDL—is converted to lysophosphatidylcholine after cleavage of a fatty acid from the sn-2 position by phospholipase A$_2$. It has
Figure 9  Schematic representation of lipid peroxidation. Polyunsaturated fatty acids are converted to lipid radicals, a process that can be inhibited by, for example, ubiquinol-10 (UQ). After molecular rearrangement, the lipid radical becomes a conjugated diene and then a peroxyl radical. This highly reactive species can attack other polyunsaturated fatty acids, thereby initiating a chain reaction. Vitamin E and ubiquinol, however, scavenge lipid peroxyl radicals, thereby breaking the chain reaction. Vitamin E can be regenerated by ascorbic acid (vitamin C) or ubiquinol-10.

It has been postulated that the released fatty acid is readily oxidized and might then become responsible for propagation of the lipid peroxidation chain reaction, as inhibitors of phospholipase A2 block the generation not only of lysophospholipids but also of lipid peroxides [31].

After peroxidation, lipid peroxides decompose and breakdown products, such as malondialdehyde (MDA) and several aldehydes, such as 4-hydroxynonenal (4-HNE), are formed. These products can react with the ε-amino groups of apoB-100, which causes an irreversible modification of the apolipoprotein, as the number of free cysteine and charged lysine residues of apoB-100 decreases. This results in reduced recognition and uptake of LDL by the LDL receptor, since the affinity of the LDL receptor is based on binding of positively charged apoB-100 to the negatively charged binding domain of the LDL receptor. This reduced uptake is compensated for by an increased affinity of these modified LDLs to the acetyl or scavenger receptors on the cell surface of macrophages. This uptake is not down-regulated and may lead to extensive lipid loading and the transformation of macrophages into foam cells (see also IV.B).

Peroxidation products are cytotoxic, and chronic irritation of endothelial cells results in lesions of the endothelial cell layer. In addition, lysophosphatidylcholine from oxidized LDLs and the expression of chemoattractant proteins like monocyte at
chemoattractant protein-1 (MCP-1), other chemoattractants, and inflammatory cytokines by damaged endothelial cells and leukocytes attract leukocytes from the circulation and initiate a local inflammation. Animal studies have shown that inflammatory cell recruitment and activation is critical for the development of atherosclerosis. For example, MCP-1 knockout mice and macrophage colony-stimulating factor (M-CSF) knockout mice are less susceptible for the development of atherosclerosis.

One of the earliest inflammatory steps in the atherogenesis is a slower rolling of leukocytes along the vascular endothelium, which proceeds by a subsequent attachment of rolling leukocytes to the vascular endothelium. In this process several adhesion molecules such as vascular cell adhesion molecule 1 (VCAM-1), intercellular adhesion molecule 1 (ICAM-1), and P-selectin play an important role. At least in vitro, these adhesion molecules are rapidly synthesized in response to several proinflammatory cytokines, such as tumor necrosis factor α (TNF-α) and interleukin-1 (IL-1). Moreover, HDL cholesterol particles can inhibit the cytokine-induced expression of adhesion molecules (VCAM-1, and E-selectin) on endothelial cells in vitro [32]. This finding may be a possible link with the antiatherogenic effect of high serum HDL cholesterol concentrations in vivo.

The presence of adhesion molecules on endothelial cells alone, however, is not enough for attachment of leukocytes to the endothelium. For this interaction leukocytes need to express ligands for these adhesion molecules. These ligands, expressed on lymphocytes and monocytes, are known as integrins. For example, very late antigen-4 (VLA-4), a β1-integrin is a ligand for VCAM-1 and lymphocyte function associated-1 (LFA-1) a β2-integrin for ICAM-1. After attachment, the proinflammatory leukocytes infiltrate into the activated endothelium, followed by a further progressing of the inflammatory response. Interestingly, blocking VLA-4 by antibodies indeed decreased leucocyte entry and fatty streak formation in mice fed an atherogenic diet [33], which shows that this integrin plays a causal role during atherosclerosis. In conclusion, this process results in a continuous recruitment of new monocytes and T lymphocytes to the place of oxidation in the endothelium and in accumulation of macrophages filled with (oxidized) LDL in the arterial intima. Also, several other processes are activated, which results in platelet aggregation, disturbance of eicosanoid homeostasis, and release of growth factors. These factors cause smooth muscle cell to proliferate and ultimately to migrate from the media to the intima. All these mechanisms together will result in the formation of fatty streaks and atherosclerotic plaques (Fig. 10).

1. Measurement of LDL Oxidation

LDL oxidation is thought to be initiated and to proceed primarily in the endothelial layer, while oxidized LDL particles are rapidly removed from the circulation. Thus, it is very difficult to quantify the LDL oxidation process in vivo, and several in vitro assays have been developed to measure LDL oxidation tendency in vitro.

a. In Vitro Copper-Mediated LDL Oxidation. Oxidative in vivo modification of LDL can be mimicked by exposure of LDL particles in vitro to redox-active metal ions, such as copper (Cu²⁺), or to reactive oxygen species, such as superoxide anion.

Esterbauer et al. [34] have developed a widely used method to determine in vitro the susceptibility of LDL to oxidation by continuous monitoring of the for-
Figure 10  Schematic representation of the formation of an atherosclerotic plaque. (1) LDL enters the intimal layer and can be oxidized by several factors, such as lipoxygenase or reactive oxygen species. (2) Oxidized LDL is cytotoxic and causes endothelial damage, (3) which results in the expression of adhesive glycoproteins to which monocytes and T lymphocytes attach. (4) The damaged endothelial cells excrete chemoattractants, which causes a continuous recruitment of monocytes and T lymphocytes. (5) These cells pass the endothelial cell layer and monocytes may become macrophages. (6) Oxidized LDL prevents return of macrophages back to the lumen and (7) the arrested macrophages absorb large amounts of oxidized LDL via the scavenger receptors and become foam cells, which may eventually lead to the formation atherosclerotic plaques.

formation of conjugated dienes, products formed after oxidation of the polyunsaturated fatty acids from the LDL particle. After LDL isolation, copper is added to the test tube to initiate the oxidation process and the formation of conjugated dienes is then quantified by measuring the change in absorbance at 234 nm (Fig. 11). This curve can be divided into three consecutive phases: the lag phase, the propagation phase, and the decomposition phase.

During the lag phase, LDL-bound lipophilic antioxidants protect the polyunsaturated fatty acids from oxidation. Tocopherols and β-carotene, for example, scavenge lipid peroxide radicals, thereby breaking the chain reaction. The antioxidant has now become a relatively stable radical, which does not induce lipid peroxidation but is also not regenerated, as in vivo may happen. Addition of the water-soluble ascorbic acid (vitamin C) that is lost during LDL isolation leads to an increase in the lag phase because this antioxidant can regenerate tocopherols. Thus, at a certain stage the LDL particle becomes depleted of antioxidants; at this point, the peroxi-
In vitro LDL oxidation. Formation of conjugated dienes during copper-catalyzed oxidation of LDL in vitro is monitored spectrophotometrically at 234 nm. The lag time before onset of rapid oxidation the maximum rate of oxidation during the propagation phase, and the maximal amount of dienes formed are used to describe LDL oxidation characteristics.

The oxidation reaction shifts to an autocatalytic process. Now the propagation phase starts during which polyunsaturated fatty acids are oxidized and converted into conjugated dienes. The absorbance at 234 nm then reaches a maximum and may eventually decrease, since the produced conjugated dienes are labile and decompose to several products, such as MDA and 4-HNE.

The oxidation profile of LDL is described by the duration of the lag time, the maximal amount of dienes formed, and the maximal rate of oxidation. A short lag time and/or a large amount of dienes is considered to reflect a high oxidative susceptibility of the LDL particle. However, interpretation of the oxidation rate is not clear.

b. Thiobarbituric Acid–Reactive Substances. LDL oxidation is also estimated by measuring in plasma the amount of thiobarbituric acid reactive substances (TBARs), such as MDA and/or MDA-like substances, which can be formed during the in vivo peroxidation process. The detection of TBARs in plasma and LDL might therefore be an indication of the possible occurrence of peroxidative injury. However, several authors regard this assay as nonspecific, as also compounds that are not a product of lipid peroxidation are measured [35].

c. Antibodies. Oxidation of LDL in vivo can be measured by the detection of autoantibodies against MDA-modified LDL in human plasma [36]. Another approach for the determination of in vivo LDL oxidation is the measurement of antibodies that cross-react with MDA-modified apoB-100, but not with native apoB-100 [37,38].

d. Isoprostanes. Isoprostanes are isomers of prostaglandins (PGs), which are produced in vivo primarily—if not exclusively—by free radical–mediated peroxidation of polyunsaturated fatty acids. Especially the F2-isoprostanes, which are isomers of the PGF$_2$$_{alpha}$, derived from peroxidation of arachidonic acid, are considered suitable markers for oxidative modifications in vivo [39]. F2-isoprostanes can be analyzed in
the circulation as well as in the urine by gas chromatography/mass spectrometry (GC-MS) or an immunoassay.

2. LDL Oxidation and Coronary Heart Disease

Although causality has not yet been proved, it would be unwise to ignore a possible role of LDL oxidation in the genesis of CHD.

Palinsky et al. [36] have demonstrated the existence of autoantibodies directed against MDA-LDL in the human circulation. Further, Ylä-Herttuala et al. [37] have shown that atherosclerotic lesions contain compounds that react with antibodies directed against MDA-modified LDL and 4-HNE-lysine, while those antibodies did not react with native LDL. In that same study it was found that LDL isolated from atherosclerotic lesions showed a substantial correspondence to the characteristics of in vitro–oxidized LDL, such as an increased amount of lysophosphatidylcholine and a chemotactic activity for monocytes. Furthermore, patients with acute myocardial infarcts or with carotid atherosclerosis show significantly higher plasma concentrations of MDA-modified LDL than control subjects [38].

Urinary immunoreactive F2-isoprostanes were higher in hypercholesterolemic patients than in controls. Moreover, urinary F2-isoprostanes were inversely related to serum LDL cholesterol concentrations and LDL vitamin E levels [40]. Also, in non-insulin-dependent diabetic patients urinary immunoreactive isoprostane levels were higher than in controls, which could be counteracted by vitamin E supplementation [41].

Not only oxidation of polyunsaturated fatty acids but also certain oxidized forms of cholesterol (oxysterols) are atherogenic and may play a role in plaque development [42]. Especially 7β-hydroxycholesterol may be a good marker for free radical–related lipid peroxidation. Two separate studies have now shown that in humans 7β-OH-cholesterol is associated with the risk for atherosclerosis [43,44]. Also feeding a mixture of various oxysterols to atherosclerosis-prone LDL receptor (−/−) and apoE (−/−) mice accelerated fatty streak lesion formation in comparison with cholesterol feeding alone [45]. For humans, however, so far no causal relation has been established between oxysteroids and lesion formation.

3. Dietary Effects

The fatty acid composition of the diet, and in particular the amount of polyunsaturated fatty acids, is reflected by the fatty acid composition of the LDL particle (Tables 4 and 5). As polyunsaturated fatty acids are more easily oxidized than monounsaturated or saturated fatty acids, it can be envisaged that LDL oxidizability is influenced by changing the dietary fatty acid composition. However, when interpreting the results it should be realized that diet always induces multiple changes in the fatty acid composition of the LDL particle. For example, the proportion of linoleic acid in the LDL particle increases after enrichment of the diet with this fatty acid, whereas the proportion of oleic acid, arachidonic acid, palmitoleic acid, and palmitic acid decreases [46].

a. Effects of Linoleate-Rich vs. Oleate-Rich Diets. Linoleic acid, an n-6 polyunsaturated fatty acid, contains more unsaturated bonds than oleic acid and is preferentially incorporated into tissue lipids. Accordingly, increasing the amount of linoleic acid in the diet at the expense of oleic acid leads to a higher proportion of
linoleic acid in the LDL particle (Table 4). Therefore, it can be expected that, at least in vitro, LDL is more easily oxidized after consumption of linoleic acid–enriched diets. Indeed, several studies have demonstrated that replacement of oleic acid in the diet for linoleic acid may result in a decreased lag time, a higher production of dienes, and a reduced oxidation rate. In addition, LDL uptake by macrophages was increased after consumption of linoleic acid–enriched diets, suggesting that in vivo the LDL was modified to a greater extent [46–50].

b. Effects of n-6 vs. n-3 Polyunsaturated Fatty Acids. Consumption of fish oils or fish oil capsules, which are rich in EPA and DHA, also affects the fatty acid composition of the LDL particle. The amount of these two n-3 polyunsaturated fatty acids increases mainly at the cost of n-6 polyunsaturated fatty acids (Table 5). Effects of fish oil relative to linoleic acid supplementation on LDL oxidation are contradictory.

Suzukawa et al. [51] found a reduction in lag time after in vitro copper-mediated LDL oxidation and an increased uptake of LDL by macrophages after supplementation with n-3 polyunsaturated fatty acids as compared with a corn oil–supplemented diet. Oostenbrug et al. [52] found dietary fish oils to increase the maximal amount of conjugated dienes formed during in vitro copper-mediated oxidation, whereas the lag time decreased. Several other studies demonstrated an increase in the amount of TBArs in plasma and in LDL after fish oil supplementation

### Table 4 Fatty Acid Composition of an LDL Particle After Consumption of a Diet Rich in Oleic Acid or Linoleic Acids [46]

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Diet (g per 100 g of fatty acid)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Oleic acid</td>
</tr>
<tr>
<td>Saturated</td>
<td>24.7</td>
</tr>
<tr>
<td>Monounsaturated</td>
<td>25.0</td>
</tr>
<tr>
<td>Polyunsaturated</td>
<td>50.3</td>
</tr>
<tr>
<td>n-6</td>
<td>45.7</td>
</tr>
<tr>
<td>n-3</td>
<td>4.1</td>
</tr>
</tbody>
</table>

### Table 5 Fatty Acid Composition of an LDL Particle After Consumption of a Diet Rich in n-6 or n-3 Polyunsaturated Fatty Acids from Corn or Fish Oil, Respectively [51]

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Diet (g per 100 g of fatty acid)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Corn oil</td>
</tr>
<tr>
<td>Saturated</td>
<td>25.5</td>
</tr>
<tr>
<td>Monounsaturated</td>
<td>23.5</td>
</tr>
<tr>
<td>Polyunsaturated</td>
<td>48.9</td>
</tr>
<tr>
<td>n-6</td>
<td>46.4</td>
</tr>
<tr>
<td>n-3</td>
<td>2.5</td>
</tr>
</tbody>
</table>
However, other studies did not demonstrate an effect of dietary fish oil relative to oils rich in linoleic acid on in vitro LDL oxidation [56,57]. Currently, there is no explanation for these contradictory results.

c. Antioxidants. Fat-soluble antioxidants, especially α-tocopherol (vitamin E), are a main protecting factor against in vitro LDL oxidation. Supplementation of vitamin E increases the tocopherol content of LDL, resulting in a higher oxidation resistance of LDL, as evidenced from an increased lag time [58]. Other antioxidants, such as carotenoids, ascorbic acid, and ubiquinol-10, also have an important impact on the oxidative resistance of LDL.

Ascorbic acid, a hydrophilic antioxidant, and ubiquinol-10, a lipophilic antioxidant, are capable of regenerating tocopherols, and addition of these antioxidants to the test tube results in a longer lag time [59–61].

Oils rich in polyunsaturated fatty acids by nature contain relatively high concentrations of antioxidants. An exception are fish oils, which have relatively low levels of fat-soluble antioxidants, but their potentially harmful effects on LDL oxidation can be counteracted by addition of vitamin E to these oils.

B. Conclusion

Several studies now strongly suggest that oxidation of LDL in vivo takes place, which results in even more harmful LDL particles. Therefore, LDL oxidation may play an important role in the formation of atherosclerotic lesions.

Replacing saturated fatty acids in the diet with linoleic acid reduces plasma LDL cholesterol concentrations. However, this dietary intervention will also increase the proportion of linoleic acid in the LDL particle thereby its in vitro oxidizability. As monounsaturated fatty acids are less readily oxidized and have comparable beneficial effects on the plasma lipoprotein profile, oleic acid might be preferred over linoleic acid. Increased in vitro oxidizability of the LDL particle might also be observed when n-3 polyunsaturated fatty acids from fish oils are added to the diet, which can be overcome by simultaneously increasing the intake of vitamin E. Although these studies clearly demonstrate that diet affects in vitro LDL oxidizability, the importance of these findings for the in vivo situation are less evident.

VIII. HEMOSTASIS

Hemostasis, derived from the Greek words for blood and standing, is a complex, delicately balanced system of interactions to keep the blood circulating as a fluid through the blood vessels. In case of imbalance, such as when a vessel is damaged, the blood stands, starts to clot, and a stable thrombus forms. This, of course, is necessary to stop a wound from bleeding. However, if a stable thrombus is formed in a small coronary artery, the artery becomes occluded, blood and oxygen supply are hampered, and a heart attack results.

The hemostatic system involves interacting processes for the formation of a stable thrombus—platelet aggregation and blood clotting—but also a mechanism to dissolve the thrombus, i.e., fibrinolysis (Fig. 12). In vivo these processes are associated and the interplay defines the prethrombotic state of the blood.

Hemostatic factors are difficult to measure. Due to the venipuncture and subsequent blood sampling and plasma preparation, platelets might become activated,
Figure 12  Processes involved in thrombus formation. The hemostatic system involves interacting processes for the formation of a stable thrombus—platelet aggregation and blood clotting—but also a mechanism to dissolve the thrombus, i.e., fibrinolysis.

which makes it very difficult to obtain a true reflection of the in vivo situation. In addition, measurements are usually made in venous fasting blood, while one is interested in thrombotic tendency in the arteries. Also, many different methods are used, which makes a comparison between studies difficult.

A. Platelet Aggregation

The activity of blood platelets is an important factor for thrombus formation. Aggregated platelets adhere to the injured blood vessel to form a hemostatic plug, excrete substances such as thrombin and calcium, and provide a phospholipid surface—all of which are important for blood coagulation.

Several mechanisms are being proposed to explain the effects of fatty acids on platelet aggregation. Differences in fatty acid composition can change the arachidonic acid content of platelet and endothelial phospholipids. Arachidonic acid acts as a substrate for thromboxane A₂ (TxA₂) in platelets and prostacycline (PGI₂) in endothelial cells, and the balance between these two eicosanoids affects platelet aggregation (Fig. 13). Fatty acids have also been reported to directly affect TxA₂ receptors on platelet membranes. A third mechanism is that differences in fatty acid composition can affect the cholesterol content of membranes, and consequently affect the fluidity of platelet membranes and platelet activation.

1. Measurement of Platelet Aggregation

A broad scale of methods is available to measure platelet aggregation in vitro. First, the blood sample needs to be anticoagulated to avoid clotting of the blood in the test tube or in the aggregometer. Different anticoagulants are used, such as citrate, which depletes the sample from calcium, and heparin or hirudin, which cause an inhibition of the conversion of prothrombin to thrombin. In vitro platelet aggregation can then be measured in whole blood, in platelet-rich plasma, or—to remove the influence of possible interfering constituents from the plasma—in a washed platelet sample. Finally, the aggregation reaction in the test tube can be triggered with many different compounds, such as collagen, ADP, arachidonic acid, and thrombin.
Figure 13  Schematic representation of platelet activation. Arachidonic acid from platelet acts as a substrate for thromboxane A2 (TxA2) and arachidonic acid from the endothelial cells for prostacycline (PGI2). The balance between these two eicosanoids affects platelet activation. TxB2 is the stable metabolite of TxA2, and 6-keto-PGF_{1a} of PGI2.

2. Platelet Aggregation and Coronary Heart Disease

In vitro platelet aggregation has been reported to be an important marker for the prediction of reoccurrence of coronary events [62]. In addition, platelet aggregation measured in whole blood was strongly associated with the prevalence of ischemic heart disease [63].

3. Dietary Fats and Platelet Aggregation

a. Total Fat Content of Diets. Renaud and coworkers [64] studied nine groups of farmers from different areas in France and Britain, who differed with respect to dietary intakes of total and saturated fatty acids. In the groups with a high consumption of total and saturated fatty acids, an increased thrombin-induced aggregation in platelet-rich plasma was observed as compared with the groups of farmers with lower intakes. In a later intervention study [65], the diets of French farmers were reduced in saturated fat content by replacing the habitually consumed dairy fat by a high linoleic acid margarine. A control group of farmers was advised not to change their diets. In the intervention group, total fat intake decreased along with the intake from saturated fatty acids, whereas the intake of dietary linoleic acid and ω-linolenic acid was increased compared with initial values. In agreement with the previous study [64], a decreased thrombin-induced aggregation was observed in the intervention group, whereas platelet aggregation did not change in the control group. However, platelet aggregation induced by ADP was significantly increased in the intervention group. From this dietary intervention, however, it was not clear whether the changes of total fat content or the changed fatty acid composition of the diets were responsible for the changes observed in platelet aggregation.

b. Dietary Fatty Acid Composition. The effects of dietary fatty acid composition were further evaluated in well-controlled dietary intervention studies, in which total fat content of the diets was kept constant and only the dietary fatty acid composition changed.

Mutanen et al. [66] observed increased ADP- and collagen-induced aggregation in platelet-rich plasma after consumption of diets rich in sunflower or rapeseed oils.
as compared with a diet rich in butterfat. In a second study, both thrombin- and ADP-induced aggregation of platelet-rich plasma was significantly higher with a high oleic acid sunflower oil diet compared with a rapeseed oil diet [67]. The authors concluded that especially the ratio of linoleic acid to α-linolenic acid of the diets was an important determinant for platelet reactivity. It was postulated that this higher ratio on diets rich in sunflower oil will result in higher concentrations of arachidonic acid, the substrate for TxA2 production, in platelet phospholipids. According to this hypothesis, oleic acid should lower platelet aggregation, as the proportion of arachidonic acid in platelet phospholipids is decreased following oleic acid consumption. However, this was not supported by the results from a small study with only three or four subjects per diet group on the effects of dietary linoleic acid compared with those of oleic acid. In this study, the oleic acid diet increased ADP- and collagen-induced platelet aggregation [68].

Effects of specific saturated fatty acids on stable metabolites of TxA2 and PGI2—TxB2 and 6-keto-PGF1α, respectively—in urine have been investigated for diets rich in stearic acid provided by cocoa butter or milk chocolate, as compared with diets rich in butterfat [69]. Although milk chocolate consumption decreased and butterfat intake increased the arachidonic acid contents of platelet phospholipids, the three diets did not change the concentrations of TxB2 and 6-keto-PGF1α in urine.

B. Coagulation

Several pathways for blood coagulation exist. The tissue factor pathway of blood coagulation, previously known as the extrinsic pathway of blood coagulation, appears to be the most important one. The factors involved in this pathway are depicted in a simplified scheme in Figure 14. Most coagulation factors are mainly present in inactivated form, except for factor VII, of which 1% circulates as activated factor VII (factor VIIa) [70]. However, Figure 14 only shows the activated coagulation factors, except for factor VII.

Thrombus formation in vivo is initiated when factor VII or factor VIIa contacts thromboplastin tissue factor, expressed, for example, after vessel injury or inflammation. A tissue factor, such as thromboplastin, is a procoagulant that is expressed only on activated endothelium. Once bound, factor VII is rapidly activated into factor VIIa. The complex of factor VIIa with tissue factor initiates a cascade of reactions, which ultimately results in the conversion of factor X into factor Xa and the generation of thrombin from prothrombin. Thrombin finally cleaves fibrinogen into fibrin, that stabilize a thrombus. However, thrombin also inhibits some coagulation factors, and the coagulation cascade is thus inhibited by one of its end products so as to prevent uncontrolled formation of fibrin.

The coagulation cascade is further regulated by the action of coagulation inhibitors. An important inhibitor of coagulation is the tissue factor pathway inhibitor (TFPI), which inhibits the activity of the tissue factor–factor VIIa complex and TFPI therefore prevents further activation of the coagulation cascade. The tissue factor–factor VIIa complex is also inhibited by antithrombin III, which also suppresses the activation of thrombin and other activated coagulation factors [71].

1. Measurement of Coagulation

Most assays measure the total amount (e.g., factor VII–antigen or fibrinogen concentrations) or the activity of circulating coagulation factors. However, these factors

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Figure 14  Schematic representation of the coagulation cascade. When factor VII contacts a tissue factor, expressed, for example, after vessel injury or inflammation, factor VII is rapidly converted to factor VIIa. The complex of factor VIIa with tissue factor initiates a cascade of reactions, which ultimately results—in the conversion prothrombin into thrombin. Thrombin cleaves fibrinogen into fibrin, which stabilizes a thrombus. Tissue factor pathway inhibitor (TFPI) and antithrombin III inhibit the coagulation cascade.

are normally present in large excess in the blood, and only a small percentage is converted to active enzymes under in vivo situations. Nowadays other assays are also available, which reflect actual in vivo coagulation. The plasma fragment 1+2 concentration reflects the amount of prothrombin actually converted to thrombin, while fibrinopeptides A and B concentrations reflect the conversion of fibrinogen to fibrin. However, in healthy subjects, concentrations of most of these markers are very low and just above detection limits, which sometimes make it difficult to implement them in dietary studies.

Many methods are available to measure factor VII. In many studies, factor VII is measured with a coagulant assay (factor VII coagulant activity). Factor VII activity, however, is measured with a two-step chromogenic assay, which depends on the rate of generation of factor Xa from factor X by factor VIIa. Both methods may give different results and do not differentiate between factor VIIa and factor VII antigen concentrations [70].

Fibrinogen concentrations are usually measured with the method of Clauss [72]. A fixed surplus of thrombin is added to diluted platelet-poor plasma samples, the clotting time recorded, and the fibrinogen concentration read from a calibrator curve.

2. Coagulation and Coronary Heart Disease

Long-term prospective epidemiological studies have consistently reported that in healthy males factor VII coagulant activity and fibrinogen concentrations were higher in subjects who developed cardiovascular diseases at a later stage of the study. Factor VII coagulant activity was particularly associated with an increased risk of dying
from cardiovascular disease [73,74]. In addition, from the Northwick Park Heart Study (NPHS) has been reported that low and, unexpectedly, also high concentrations of antithrombin III were associated with increased deaths from CHD [75]. It must be noted that these prospective studies have only been carried out in males; whether associations for females are similar awaits confirmation.

3. Dietary Fats and Coagulation

a. Total Fat Content of Diets. Marckmann and colleagues [76] have investigated both short and longer term effects of low-fat/high-fiber diets on human blood coagulation. In an 8-month study it was found a low-fat/high-fiber diet significantly decreased plasma factor VII coagulant activity by 5–10%, but only in the first 2 months and in the last months of dietary intervention. Absence of effects in the middle-study period was explained by the fact that the subjects in this period did not follow the dietary guidelines strictly due to allowed study holidays. Plasma fibrinogen concentrations were not changed [76]. The results of this study were confirmed in a study of shorter duration [77].

In another study, Marckmann et al. [78] investigated whether the low-fat or the high-fiber component of diets was responsible for decreased factor VII coagulant activity found in earlier studies. The experimental diets of this trial only differed in their fat content (39 vs. 31% of energy) and carbohydrate content (47 vs. 54% of energy). Factor VII coagulant activity was similar on the low-fat and high-fat diets. In addition, no changes in fibrinogen concentrations were found. This limited number of studies indicates reducing effects on factor VII coagulation activity of low-fat/high-fiber diets. However, more controlled studies are needed to definitely address whether the reduced fat content of low-fat diets, the increased fiber content, or a combination of these two dietary factors are responsible for the decreased factor VII coagulant activity of such diets.

b. Dietary Fatty Acid Composition. The effects of saturated compared with unsaturated fatty acids have been studied in several experiments. Factor VII antigen, factor VII coagulant activity, and estimated factor VIIa increased with diets high in saturated fatty acids compared with unsaturated fatty acids [79]. This study included only four men and one woman, and the experimental diets were extremely high (62% of energy) in fat content. Additional experiments were designed to investigate the effects of different ratios of polyunsaturated to saturated fatty acid content (P/S ratio), at more realistic fat intakes of 32% of energy [80] or 40% of energy [81]. One study [80] did not indicate changes of factor VII coagulant activity on a diet with a P/S ratio of 0.3 versus a diet with a P/S ratio of 0.9. Another study [81] reported only slightly increased (1.7%) factor VII antigen concentrations when the dietary P/S ratio was decreased from 3.0 to 0.3. Fibrinogen concentrations were not measured in these studies.

Diets enriched in n-3 fatty acids from fish did not change factor VII coagulant activity [82,83] or fibrinogen concentrations [82–84].

A study of Almendingen et al. [85] investigated the effects of trans fatty acids in diets enriched in hydrogenated fish oil or hydrogenated soybean oil compared with a diet enriched in butterfat. The diet enriched in butterfat showed slightly increased fibrinogen concentrations as compared with the hydrogenated fish oil diet. No significant differences in the levels of factor VII or fibrin degradation products were observed.
Recently, well-controlled studies investigating effects of specific saturated fatty acids have been published. Tholstrup et al. [86] reported that diets rich in lauric plus myristic acids or palmitic acid increased factor VII coagulant activity as compared with a diet rich in stearic acid. It was also suggested that the turnover of prothrombin was increased on the lauric plus myristic acids diet as fragment 1+2 concentrations were higher on such a diet than on a stearic acid diet [87] (Fig. 15). However, stearic acid increased fibrinogen concentrations compared with the lauric plus myristic acid diet [87]. In a second study, Tholstrup et al. [13] investigated diets rich in myristic acid or palmitic acid. On the myristic acid diet, subjects showed an increase in factor VII coagulant activity. Thus, these studies [13,86] suggest that saturated fatty acids, except for stearic acid and probably MCFA, increase factor VII coagulant activity. However, stearic acid may have an unfavorable effect on fibrinogen concentrations.

C. Fibrinolysis

The process involved in thrombus dissolution, and thus the conversion of fibrin into fibrin degradation products, is called fibrinolysis. A simplified scheme of the fibrinolytic pathway is given in Figure 16. The central reaction in the fibrinolytic process is the conversion of plasminogen into plasmin, which is regulated by the action of tissue plasminogen activator (t-PA). The fibrinolytic capacity of blood is regulated by inhibiting t-PA activity by the action of plasminogen activator inhibitor type-1 (PAI-1), while plasmin is inhibited mainly by α2-antiplasmin [88].

1. Measurement of Fibrinolytic Capacity of Plasma

The fibrinolytic capacity of plasma can be measured by global tests or more specific assays. The global tests include the dilute clot lysis time, the euglobulin clot lysis
time, and the fibrin plate assay. The dilute and euglobulin clot lysis time measure total fibrinolytic capacity. The total blood sample or the insoluble protein sample (euglobulin fraction) is diluted with a buffer, clotted, and the lysis time of the clot recorded. In the fibrin plate assay, a standard volume of the euglobulin fraction of plasma is added to standardized plasminogen-rich fibrin plates and the amount of lysis is recorded. In more specific assays, total plasma concentrations of t-PA and PAI-1 can be determined with enzyme-linked immunosorbent assays. Plasma t-PA and PAI-1 activities can be estimated with a chromogenic assay.

2. Fibrinolysis and Coronary Heart Disease

From the NPHS has been reported that low fibrinolytic capacity of plasma, measured as clot lysis time, was significantly associated with increased CHD risk in men aged 40–54 [89]. In addition, Hamsten et al. [90] have reported that higher concentrations of PAI-1 were associated with increased risk of reoccurrence of coronary events.

3. Dietary Fats and Fibrinolysis

a. Total Fat Content of Diets. Marckmann et al. [76,77] found in both a short term (2 week) and a longer term trial (8 months) that plasma euglobulin fibrinolytic capacity and plasma t-PA activity were increased on low-fat/high fiber diets as compared with high-fat/low-fiber diets (Fig. 17). No changes were observed in t-PA and PAI-1 antigen concentrations. In agreement with these results, Mehrabian et al. [91] reported decreased plasminogen and PAI-1 activities, after consumption of a low-fat/high-fiber/low-cholesterol diet with less than 10% of energy from fat for 21 days. This was associated with decreased t-PA antigen concentrations. However, from these studies it is not possible to conclude whether the changes observed were due to the lower fat or cholesterol contents, or the higher fiber content of the diets.

In another study, Marckmann et al. [78] have also compared two diets, which differed in total fat content though fiber intake and the relative fatty acid composition...
were comparable. Their results indicated that plasma euglobulin fibrinolytic capacity on a high-fat diet (39% of energy from fat) did not change compared with a low-fat diet (31% of energy). It was concluded that an isolated reduction of total fat content of the diet does not affect the fibrinolytic capacity of the blood but that concomitant changes in the diet, as for example changes in fatty acid composition or fiber content, are necessary to provoke changes of fibrinolytic factors.

b. Dietary Fatty Acid Composition. Effects of different fatty acid composition on fibrinolytic capacity have been investigated extensively for fatty acids from fish, although recently additional data have become available for the effects of other fatty acids.

High intakes of fish may affect fibrinolytic capacity. Brown and Roberts [83] studied the effects of a daily consumption of 200 g of lean fish, with or without fish oil supplement. They observed an apparent enhancement of plasma euglobin fibrinolytic capacity, which tended to be accentuated with fish oil supplementation. However, compared with meat diets, diets enriched with fatty fish have also been associated with an unfavorable increase in PAI-1 activity [82,84]. Increased PAI-1 activity has also been reported on diets enriched in partially hydrogenated soybean oil compared with a butterfat diet [85].

Tholsrup and coworkers [86] did not report unfavorable effects of lauric plus myristic acids or palmitic acid diets relative to stearic acid on euglobulin fibrinolytic capacity, t-PA or PAI-1 activity. Also myristic acid and palmitic acid diets did not change these variables as compared with habitual diets [13].

D. Conclusion

Results of platelet aggregation studies are difficult to interpret because of large differences in the methods used. Lowering dietary fat intake decreases in vitro platelet
aggregation tendency induced by some antagonists but increases in vitro platelet aggregation by other antagonists. Also, the effects of dietary fatty acid composition on in vitro platelet aggregation are contradictory.

To affect coagulation and fibrinolytic factors, it appears that lowering fat intake must be accompanied by changes in dietary fatty acid composition and/or increased fiber intakes. Favorable effects on blood coagulation can be obtained by lowering the content of the major dietary saturated fatty acids and replacing them with unsaturated fatty acids.

REFERENCES

50. K. D. Croft, P. Williams, S. Dimmott, R. Abu-Amsha, and L. J. Beilin. Oxidation of low density lipoproteins: Effect of antioxidant content, fatty acid composition and intrinsic...


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Conjugated Linoleic Acids: Nutrition and Biology

BRUCE A. WATKINS and YONG LI

Purdue University, West Lafayette, Indiana

I. INTRODUCTION

Conjugated linoleic acids (CLAs) are a family of positional and geometric isomers of octadecadienoic acid (18:2). Double bonds in CLA are conjugated and not separated by a methylene group (—CH₂—) as in linoleic acid (LA or 18:2n-6), an ω-6 essential fatty acid. The CLA isomers are found in many foods (1) but are predominant in products derived from ruminant sources (beef, lamb, and dairy) because of the process of bacterial biohydrogenation of polyunsaturated fatty acids (PUFAs) in the rumen (1–3).

The highest concentrations of CLA in food are present in dairy products (4,5) and fat in the meats of lamb, veal calves, and cattle (6). In most cases, the cis-9, trans-11 isomer is the chief isomer of CLA found in food. Estimates of CLA intake range from 0.3 to 1.5 g/person/day and appear to be dependent on gender and the intake of food from animal and vegetable origins (4).

Since the finding that CLAs isolated from grilled beef inhibited chemically induced cancer (7–9), numerous studies have been initiated to investigate the physical, biochemical, and physiological properties of CLA isomers. The growing body of literature on CLA suggests that these isomeric conjugated fatty acids possess potent biological activities that may benefit human health.

II. CHEMISTRY

The acronym CLA is used to describe a family of octadecadienoic acid (18:2) isomers that possess a pair of conjugated double bonds along the alkyl chain from carbon 2
Octadecadienoic acids have been reported to contain conjugated double bonds at positions 7,9; 8,10; 9,11; 10,12; 11,13; and 12,14 (counting from the carboxyl end of the molecule) in chemically prepared CLA mixtures or natural products (10–13). The positional conjugated diene isomers can occur in one or more of the following four geometric configurations: cis,trans; trans,cis; cis,cis; or trans,trans; which would give 24 possible isomers of CLA (10). Many of these isomers were found in commercially available preparations of CLA produced under alkaline conditions from vegetable oils containing a high concentration of LA (14).

The most common CLA isomer found in meat from ruminant species and bovine dairy food products is octadeca-c9,t11-dienoic acid (15), even though minor components, such as the t7,c9, t8,c10, t10,c12, t11,c13, c11,t13, and t12,t14 isomers, and their cis,cis, trans,trans isomers were also reported in these products (11,12). Two trivial names have been proposed for the c9,t11 isomer: bovinic acid (16) and rumenic acid (17). The name bovinic acid is considered to be too restrictive for CLA because the c9,t11 isomer is not only produced in the rumen of the bovine but also produced by other ruminant animals by the same mechanism.

The CLA in ruminant meat and dairy products is believed to be formed by bacterial isomerization of LA and possibly α-linolenic acid (18:3n-3) from grains and forages to the c9,t11-18:2 in the rumen of these animals (1,2,18). CLA may also be formed during cooking and processing of foods (18). Presently CLA mixtures and pure isomers are available from various commercial sources. The composition of CLA products should be carefully checked before they are used in research work since the isomeric distribution varies largely between manufacturers and even between batches made by the same manufacturer (19,20).

### III. ANALYSIS OF CLA IN FOOD AND BIOLOGICAL SAMPLES

As research continues on the actions of CLA, it is critical to have accurate compositional analyses of food and biological samples to correctly interpret data from investigations. The analysis of CLA in food and other biological samples follows the general guidelines for the determination of fatty acid composition with particular emphasis on the derivatization methods. The conjugated double bonds in CLA are less stable than the methylene-interrupted double bonds in LA. It is reported that the stability of CLA is similar to that of arachidonic acid (20:4n-6) and docosahexaenoic acid (DHA, 22:6n-3) (21,22). Therefore, preventing isomerization and oxidation of these labile fatty acids is critical for a successful analysis.

The most common analyzed form of fatty acids is the fatty acid methyl ester (FAME). Both acid and alkaline catalysts can be used to prepare FAME for gas chromatography (GC) analysis. However, since acid catalysts, such as BF₃, will change the double-bond configuration in conjugated dienes and generate artifact CLA isomers (23–25), their use is not recommended in the analysis of CLA.

The alkaline catalysts perform best for lipids containing fatty acids with unique conjugated diene structures. Isomerization and artifacts are not produced when sodium methoxide or tetramethylguanidine (TMG) are used as transesterification agents; however, they do not methylate free fatty acids and N-linked (amide bond) fatty acids as those found in sphingolipids. Therefore, these methods are not suitable for samples with high acid values (high free fatty acid content). There is no single
method that works optimally in all situations. The researcher needs to know the nature of the sample and select a suitable method accordingly.

Various instruments are adopted to separate and characterize the CLA isomers. A gas chromatograph equipped with a polar 30- to 100-m capillary column and a flame ionization detector is the most widely used instrument to characterize CLA isomers. Generally, GC columns have a limited capacity to fully separate all geometric and positional isomers of CLA; therefore, it is impossible to identify every individual CLA isomer by conventional capillary GC analysis. Mossoba et al. (26) reported that 10 CLA peaks were resolved in a commercial CLA preparation with a gas chromatograph equipped with a 100-m column (SP 2560, Supelco Inc., Bellefonte, PA or CP-Sil 88, Chrompack, Bridgewater, NJ). In the aforementioned procedure the 10 peaks resolved may actually represent more than 15 CLA isomers. A more powerful approach has been the application of argentation (silver ion) high-performance liquid chromatography (Ag⁺-HPLC) using a UV detector at a wavelength of 233 nm. Sehat et al. (10) described such a method that resolves CLA isomers according to geometric configuration and position of the conjugated diene structure. Moreover, Mossoba et al. (27) reported that 16 isomers of a CLA sample were resolved by an Ag⁺-HPLC method. Although the Ag⁺-HPLC method is more powerful than GC in resolving CLA isomers, it cannot be used to quantify other fatty acids due to the limitation of UV detectors and HPLC columns. Therefore, ideally the two methods should be combined for CLA analysis, using GC for the general fatty acid analysis and Ag⁺-HPLC for the conjugated dienes. The results of these two procedures would provide a more complete profile of CLA isomers in food and biological samples.

IV. CLA CONTENT IN FOOD PRODUCTS AND BIOLOGICAL SAMPLES

Cheese is a chief source of dietary CLA in animal-derived products. The CLA concentrations in various dairy products (cheeses, milk, butter, buttermilk, sour cream, ice cream, and yogurt) range from 0.55 to 24 mg/g fat. The average CLA content in milk is about 10 mg/g milk fat (4,28). The largest variation in the amount of CLA is found in various natural cheeses ranging from 0.55 to 24 mg/g of fat. Seven CLA peaks that could represent nine isomers were present in dairy products; among these c9,t11, t10,c12, t9,t11, and t10,t12 accounted for more than 89% (18). The CLA content in cheeses is primarily dependent on the CLA content in the milk, which varies in CLA concentration due to seasonal variation, geography, nutrition of the cow, and management practices. In addition, CLA content of cheese, to a limited extent, is affected by the production process and maturation (29).

Reported values for CLA content in beef muscle vary considerably from 1.2 to 9.9 mg/g fat (1,18,30,31). Fats and meats from ruminant species are a rich natural source of CLA, and the reported values ranged from 2.7 to 5.6 mg CLA/g fat in lamb, veal, and beef. Fritsche and Fritsche (32) reported that the amount of the c9,t11, 18:2 isomer in beef averaged 0.76% of total FAME for fat samples from bulls and 0.86% for fat from steers. Minor isomers, e.g., t9,c11, c9,c11, and t9,t11, were also found in beef fat samples. Others have reported that the c9,t11 18:2 content in beef ranged from 1.7 to 6.5 mg/g fat (30) and 0.65% of total FAME in beef fillet (4).
CLA is also present in small amounts in other food products. Turkey meat has the highest CLA content of 2.5 mg/g fat for nonruminant species (18). Chicken contained CLA (0.9 mg/g fat) as did pork (0.6 mg/g fat) with c9,t11 being the major isomer (84% and 82%, respectively) (1). The amount of CLA in chicken egg yolk lipids ranges from 0 to 0.6 mg/g of fat (13,3–36). CLA was found in plant oils (0.1–0.7 mg/g fat) and selected seafood (0.3–0.6 mg/g fat) in small quantities (1). Unlike in ruminant derived food where c9,t11 is the chief isomer of CLA, this isomer accounts for only 38–47% of the total CLA in plant oils and rather interestingly appears to be absent in seafood lipids. Banni et al. (37) carried out a series of analyses to characterize the fatty acids with conjugated dienes in partially hydrogenated oil (mixture of partially hydrogenated soybean oil and palm oil) and confirmed the presence of CLA isomers in these oils. Moreover, Mossoba and coinvestigators (26) reported that conjugated cis,trans and trans,trans 18:2 isomers were present in hydrogenated soybean oil and margarine.

CLA has been identified in various human tissues, such as adipose, serum, breast milk, and in bile and duodenal secretions (38). Fogerty et al. (39) reported 5.8 mg/g fat of CLA in human milk for subjects consuming a normal Australian diet. Precht and Molkentin (40) reported a value of 3.8 mg/g fat (range 2.2–6.0 mg/g fat) in human milk obtained from 40 German women. McGuire et al. (16) analyzed 14 human milk samples from subjects in the Pacific Northwest and reported that CLA values ranged from 2.23 to 5.43 mg/g fat (mean 3.81 mg/g fat) or from 0.02 to 0.30 mg/g on a milk weight basis. In this last study, all milk samples contained 83–100% of the c9,t11 isomer, and in 8 of the 14 samples the c9,t11 isomer was the only form observed. Jensen et al. (41) reported a lower level of CLA in human milk that ranged from 1.4 to 2.8 mg/g fat with an average of 1.8 mg/g fat. Based on these data, CLA in human milk for Western societies ranged from 1.4 to 5.8 mg/g fat.

The concentration of CLA found in human plasma and serum appears to be linked to dietary fat type and food consumption patterns. Herbel et al. (38) reported that plasma CLA concentration ranged from 6.4 to 7.3 μmol/L in human subjects that were given a high level of safflower oil for 6 weeks. The high LA intake from safflower oil did not increase the plasma level of CLA, indicating that dietary LA is not converted to CLA in these subjects. In another study, the effect of trans fatty acid intake on serum CLA was examined (42). Eighty human subjects were put on a diet high in saturated fat mainly from dairy for 5 weeks. Then they were separated into two dietary groups: 40 subjects were assigned to a diet high in trans fatty acids from partially hydrogenated vegetable oil, and the other 40 given a similar diet high in stearic acid. At the termination of the study, serum samples from the trans diet group (CLA was 0.43% of total fatty acids) contained 30% more CLA than samples obtained while on the dairy fat diet (0.32% CLA). Samples from those given the stearic acid diet had only half of the amount of CLA compared to those given the dairy fat diet. These data indicate that a possible relationship exists between trans fatty acid intake and serum CLA concentration. In another dietary intervention study of nine healthy men, Cheddar cheese was added to their diet at a level of 112 g/day for 4 weeks. Plasma CLA concentration increased from 7.1 to 9.6 μmol/L at the end of the cheese supplement period and was maintained at 7.8 μmol/L after another 4 weeks following intervention (43).

The CLA concentration and isomeric distribution have recently been characterized in human adipose tissue. Fritsche et al. (44) reported that human subcutaneous
adipose tissue contained two major CLA isomers, both c9,t11 and t9,t11, and two
minor isomers t9,c11 and c9,c11. The presence of CLA in human tissues is pri-

V. NUTRITION AND BIOLOGY OF CLA

CLA is the only known anticarcinogen tested in animals associated with foods orig-
inating from animal sources. Ha and coworkers (9) provided one of the earliest obser-
vations that CLA from beef was protective against chemically induced cancer. In that study, CLA isolated from extracts of grilled ground beef were found to reduce skin tumors in mice treated with 7,12-dimethylbenz[a]anthracene (DMBA), a known carcino-
gen (9). Since then, numerous researchers have reported the effects of CLA iso-
ers. The research on CLA has relied entirely on animal models and cell culture systems employing isomeric mixtures of CLA. Recent studies in human subjects have been negative (20,51,52). The purported properties of CLA include anticarcin-
genic (9,53–59,60–62) and antiatherosclerotic (63,64). Other CLA effects include 
antioxidative (54,55,63,64) and immunomodulative (47,65–68).

More recently, preliminary data suggest that CLA may have a role in control-
ling obesity (69–71), reducing the risk of diabetes (72), and modulating bone me-
tabolism (73,74). In addition, some studies indicate that the biological effects of CLA 
are modulated by dietary sources of long chain n-3 fatty acids (68,73,74).

Considerable research has been done to examine the anticarcinogenic properties 
of CLA. Isomers of CLA have been shown to reduce chemically induced tumor-
genesis in rat mammary gland and colon (53,55–58, 60,62,75). Moreover, CLA 
modulated chemically induced carcinogenesis in mouse skin (9,76) and forestomach 
(54). Sources of CLA also inhibited the growth of human tumor cell lines in culture

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and in SCID (severe combined immunodeficiency) mice (59,79). Recent investigations with CLA failed to show a benefit on rodent models of colon carcinogenesis (80) and prostate metastasis (81).

A. Cancer

In dietary preparations, CLA provided a potent anticarcinogenic effect in a rat mammary cancer model. Ip et al. (55) found that administering weanling rats CLA at 0.5–1.5% of the diet for 2 weeks before DMBA induction (oral intubation of 10 mg in 1 mL corn oil) and continuing for 36 weeks resulted in a significant reduction in tumor incidence. The effect of CLA was maximized at a dietary level of 1% in this study. In another experiment, much lower doses of dietary CLA (0.05–0.5%) were given to rats (53). The dietary treatments induced a dose-dependent inhibition in mammary tumor yield with a lower dose of 5 mg of DMBA. In the 5-week short-term feeding experiment of 1% dietary CLA (53), rats were given CLA for 5 weeks and the carcinogens were introduced 1 week before the end of CLA treatment. Total tumor yield was reduced by 39% and 34% in the DMBA and methylnitrosourea (MNU) induction groups with CLA treatment, respectively, at the end of the experiment. The short-term feeding period corresponded to the maturation of the rat mammary gland to adult stage morphology, and the supplement of CLA was stopped shortly after the induction with the carcinogens. Therefore, these results suggest that CLA may have a direct modulating effect on susceptibility of the target organ to neoplastic transformation.

In a subsequent dietary study using the rat DMBA model, the inhibitory effect of CLA on induced mammary cancer was found to be independent of fat type (n-3 fatty acids were investigated in the study) provided in the diet (56). When providing rats with a CLA supplement, the fatty acid isomers were incorporated into the mammary gland lipids, and the amount in the neutral lipids greatly exceeded that in phospholipids (57,82). When CLA was removed from the diet after 4 and 8 weeks of feeding, neutral lipid- and phospholipid-CLA returned to basal values in about 4 and 8 weeks, respectively. The author observed that the rate of CLA disappearance in neutral lipids subsequent to CLA withdrawal closely paralleled the occurrence of new tumors in the rat target tissue. It appears that the tissue accumulation of CLA is important to its antitumor effect and that the concentration of CLA in neutral lipid may be a more sensitive biomarker of tumor protection than the amount in phospholipid for this model. The effect of CLA on cancer was maximized at a dietary level of 1% as shown in several studies, and increasing the dietary level of CLA did not afford additional protection (55,82).

The antitumor effect of naturally produced CLA was tested over a chemically prepared CLA mixture from LA which was used in previous studies (83). During the study, rats were given a diet with a high CLA content butter fat (equivalent to 0.8% CLA in the diet) and a low-level CLA diet that served as the control group (equivalent to 0.1% CLA in the diet) for 1 month before tumor induction using MNU. After tumor induction, rats were changed to a regular diet without CLA for 24 weeks before the termination of the experiment. CLA in butter inhibited the tumor yield by 53%. CLA was also shown to reduce the population of mammary terminal end bud (TEB) cells, the cells that are the primary target of attack by carcinogens, and the proliferation of TEB cells. Similar effects reported by Banni et al. (62) revealed that
providing rats with 0.5% and 1% CLA in the diet produced a graded and parallel reduction in TEB density and mammary tumor yield, and no further decrease was observed in either parameter in rats given 1.5% and 2% CLA. These studies support the hypothesis that exposure to CLA during the time of mammary gland maturation may modify the developmental potential of the target cells that are normally susceptible to carcinogen-induced transformation. In contrast to the previous experiments using a chemically prepared CLA mixture, the latter experiments, for the first time, showed that the natural form of CLA found in dairy foods is active in reducing the incidence of mammary tumors in rats.

The anticarcinogenic properties of CLA isomers were tested in various human cell culture models. Cunningham et al. (61) treated human normal mammary cells and MCF-7 breast cancer cells with LA, CLA, and eicosanoid synthesis inhibitors in cell culture. Fatty acids were complexed with bovine serum albumin (BSA) prior to introduction of treatments into the culture media at concentrations of 0–3.57 × 10^5 M. The results showed that CLA inhibited thymidine incorporation in normal and MCF-7 cancer cells. The treatment consisting of CLA plus nordihydroguaiaretic acid (NDGA), a lipoxygenase inhibitor, suppressed the growth of MCF-7 cancer cells synergistically. The authors suggested that the growth suppression was augmented by CLA through inhibition of leukotriene synthesis.

The effects of CLA on the growth of three different lung adenocarcinoma cell lines (A-427, SK-LU-1, A549) and one human glioblastoma cell line (A-172) were examined by Schonberg and Krokan (77). There was a dose-dependent reduction in proliferation of the lung adenocarcinoma cell lines with A-427 being the most sensitive, but CLA had virtually no effect on the A-172 cell line. In addition, LA had no inhibitory effect on either of these cell lines. A significant increase in lipid peroxidation (measured by the formation of malondialdehyde, MDA) was observed after exposure of the lung adenocarcinoma cell lines to 40 μM CLA. The level of MDA was approximately twofold higher than after exposure to 40 μM LA. Treatment with vitamin E (30 μM) totally abolished the formation of MDA; however, cell growth rates were only partially restored. These data might suggest that cytotoxic lipid peroxidation products are only in part responsible for the growth inhibitory effects of CLA. Further research is needed to elucidate a possible mechanism of CLA action in lipid peroxidation and disease.

The anticarcinogenic effect of CLA on a human prostatic cancer cell line was investigated in animal models. Cesano et al. (59) tested the effect of both CLA and LA against the human prostatic cancer cell line DU-145 in a mouse model. Mice were given a control diet and either LA or CLA at 1% of diet for 2 weeks before the subcutaneous inoculation of cancer cells. Mice were then maintained on the same dietary treatments for 10 more weeks before the termination of the study. Mice given the CLA-supplemented diet displayed not only smaller local tumors than those given the control diet, but also a drastic reduction in lung metastases. Mice given the LA diet had increased local tumor loads as compared with the CLA-treated and control groups. The aforementioned studies suggest that CLA and LA have distinct biological effects.

The anticarcinogenic properties of CLA isomers might be limited to certain types of cancers and may not be effective under some experimental conditions (including dietary sources of phytochemicals and nutraceutical fatty acids such as the n-3 PUFAs) (84). Wong et al. (85) showed that providing mice diets containing from
0.1% to 0.9% CLA for 2 weeks before infusing with WAZ-2T metastatic mammary tumor cells into the right inguinal mammary gland did not affect mammary tumor latency, tumor incidence, or volume. Therefore, dietary CLA might be less effective on the growth of an established and aggressive mammary tumor.

The anticarcinogenic effects of CLA isomers have been exclusively tested in animal models and cell culture systems. There is no direct evidence that these fatty acids protect against carcinogenesis in the human. The most promising evidence to date that dietary sources of CLA are beneficial is from epidemiological studies that link milk consumption to reduced breast cancer. In one recent study, Knekt et al. (86) found a significant inverse relationship between milk intake and breast cancer incidence among 4697 initially cancer-free Finish women over a 25-year follow-up period. It was found that the risk of breast cancer was halved in women who consumed more than 620 mL milk per day compared with those consuming less than 370 mL/day and suggested that CLA was the active component. The average amount of CLA in milk is approximately 10 mg/g of milk fat. Based on the average fat content of 35 g/L of milk (87), the daily intake of CLA for these women could be 217 mg for those consuming 620 mL milk per day and 130 mg for those consuming 370 mL milk per day. However, the calculated CLA intake level for these women is still far below the estimated CLA level of 1 g/day that would be significant for cancer prevention (53). The results from the epidemiological studies and reduced incidence of breast cancer could suggest that dairy fats contain other factors that reduce cancer risk or other lifestyle and dietary factors associated with these populations, e.g., fish intake (n-3 fatty acids), that would influence cancer incidence. Experiments designed for investigating the role of CLA in human cancer are needed to identify the relationships between these fatty acids and other lifestyle factors contributing to lower cancer risk.

B. Cardiovascular Disease

The potential health benefits of CLA may extend beyond cancer to the prevention of congestive cardiovascular diseases since these fatty acid isomers were shown to reduce atherogenesis in animal studies (63,64). Lee et al. (63) assessed the effect of CLA on atherosclerosis in rabbits by supplementing 0.5 g per animal per day for 22 weeks. In comparison with control rabbits that were given a similar diet containing 14% fat and 0.1% cholesterol without CLA, total and LDL cholesterol and triacylglycerol (TAG) levels in blood were markedly lower in those given CLA. At the same time, the ratio of LDL cholesterol to HDL cholesterol and ratio of total cholesterol to HDL cholesterol were significantly reduced in the group given CLA. Rabbits given CLA also showed less atherosclerosis in the aorta. In another study by Nicolosi et al. (64), CLA effects on plasma lipoproteins and aortic atherosclerosis were examined. Fifty hamsters were divided into five groups of ten and given 0 (control), 0.06 (low), 0.11 (medium), and 1.1 (high) percentage of total dietary energy of CLA or 1.1% of total energy as LA. Animals given the CLA-containing diets collectively had significantly reduced levels of plasma total cholesterol, non-HDL cholesterol (combined VLDL and LDL) and triacylglycerols with no effect on HDL cholesterol, in comparison with the controls. For the CLA treatment, a tocopherol sparing effect was observed compared to the control group. Morphometric analysis of aortas revealed less early atherosclerosis in the CLA-treated hamsters compared
with the control group. Interestingly, the LA treatment showed a similar effect compared with CLA in this study. Yeung et al. (88) reported that by feeding hamsters 20 g/kg dietary linoleic acid or CLA, serum total fasting cholesterol and triacylglycerol were significantly reduced compared with the control which had no fatty acid supplementation. However, the cholesterol-lowering mechanism for LA and CLA appeared to be different. CLA lowered the activity of intestinal acyl CoA–cholesterol acyltransferase (ACAT) while LA had no effect on this enzyme, indicating that CLA could affect the absorption of dietary cholesterol. In rats fed (up to 1.5% dietary CLA for 60 days) the ratio of HDL cholesterol/TC and serum TAG concentrations were significantly elevated (89). These studies showed that CLA was hypocholesterolemic and antiatherogenic; however, a recent experiment using C57BL/6 mice given atherogenic diets with added CLA (0.5% and 0.25% of the diet) demonstrated increased development of aortic fatty streaks despite a change in serum lipoprotein profiles that could be considered less atherogenic (90).

There is currently no conclusive evidence that CLA protects against early atherogenesis. It was initially proposed that CLA was able to protect LDL particles from oxidation (54,55,63,64); however, a recent in vitro investigation by ven den Berg et al. (21) showed no antioxidant properties attributable to CLA. Also, CLA showed its positive effect on atherosclerosis in rabbits but not in mice, which could indicate a species difference for the response to dietary CLA supplementation.

Benito et al. (52) showed that although CLA isomers were incorporated into platelet lipids, antithrombotic properties, such as blood clotting parameters and in vitro platelet aggregation, were not affected by daily supplement of 3.9 g CLA to a typical Western diet for 63 days in healthy adult females.

In a human study involving 17 female subjects, CLA was given as a dietary supplement at 3.9 g/day for 93 days. Blood cholesterol or lipoprotein levels were not altered by CLA supplementation in healthy and normolipidemic subjects. In this investigation the short-term supplementation with CLA did not afford health benefits for the prevention of atherosclerosis (20).

C. Body Fat and Lipid Metabolism

Numerous studies in growing animals demonstrated that CLA reduced fat deposition and increased lean body mass (69–71, 91–94). When CLA (0.5–1% of diet) was given to AKR/J male mice (39 days old), it produced a rapid and marked decrease in fat accumulation and an increase in protein accumulation without any major effects on food intake (69). Park et al. (93) provided diets with a lower dietary CLA level (5% corn oil + 0.5% CLA) to 6-week-old ICR mice and observed a significantly reduced body fat content by 57% (male) and 60% (female), and increased lean body mass relative to control mice that were given a diet containing 5.5% corn oil. Similar effects of CLA on body fat accumulation were shown in rats. Yamasaki and colleagues (70) studied the effect of CLA on liver and different adipose tissues in 4-week-old male rats given diets (AIN-93G with 7% safflower oil as a control) containing 1% and 2% dietary CLA (at the expense of the safflower oil) for 3 weeks. They observed reduced levels of TAGs and nonesterified fatty acids in the liver and white adipose tissue without significant changes in lipids of brown adipose tissue. In another study, Yamasaki et al. (95) reported that dietary CLA treatments (0%, 1%, and 2% of the diet) dose-dependently accelerated the release of lipids in white adi-
pose tissue and increased the clearance rate of serum nonesterified fatty acids. In addition, CLA treatment induced a nonsignificant increase in liver size and liver TBAR (thiobarbituric acid reactive substances) levels were significant higher in rats given the 2% CLA diet indicating a morphological change in liver caused by increased peroxidation.

The effect of CLA on body composition was also investigated in a randomized, double-blind, placebo-controlled study including 60 overweight or obese volunteers with body mass index (BMI) ranging from 27.5–39.0 kg/m² (96). The subjects were divided into two groups receiving 3.4 g CLA or placebo (4.5 g olive oil) daily for 12 weeks. No difference in adverse events or other safety parameters was found between the treatment groups. Small changes in the laboratory safety data were not regarded as clinically significant. In the CLA group, mean weight was reduced by 1.1 kg (paired t-test, \( p = 0.005 \)), while mean BMI was reduced by 0.4 kg/m² (\( p = 0.007 \)). However the overall treatment of CLA on body weight and BMI was not significant. The results indicate that CLA in the given dose did not adversely impact a healthy population based on the safety parameters investigated. In another study, Blankson et al. (97) reported that feeding overweight or obese human subjects (BMI 25–35 kg/m²) up to 6.8 g of CLA per day for 12 weeks reduced the total body mass. The CLA effect on body fat mass reduction peaked at 3.4 g/day and higher dietary intake did not show a further benefit.

Repartitioning of fat to lean was reported in growing pigs fed CLA. Dugan et al. (91) demonstrated that in pigs (male and female) given a cereal-based basal diet containing either 2% CLA compared to sunflower oil led to reduced subcutaneous fat deposition and increased lean body mass. Pigs provided with CLA also had reduced feed intake (5.2%) and increased feed efficiency (5.9%) compared with pigs fed sunflower oil. In another study, finisher pigs were offered six treatments having from 0 to 10 g CLA/kg of diet for 8 weeks (92). Dietary CLA treatments resulted in increased feed efficiency and lean tissue deposition, and decreased fat deposition (decreased by 31% at the highest CLA level) in growing pigs.

The fat partitioning effect of CLA was further examined in the adipocyte 3T3-L1 cell culture. Park et al. (93) found that when added during fat accretion in the 3T3-L1 adipocyte culture, CLA (1 × 10^4 M complexed with albumin) reduced lipoprotein lipase activity and enhanced lipolysis leading to less fat deposition. In addition, skeletal muscle from mice fed CLA exhibited elevated carnitine palmitoyltransferase activity, which indicates elevated \( \beta \) oxidation of fatty acids. In a recent study using the same cell culture system (71), a specific CLA isomer, t10,c12-18:2, was found to reduce lipoprotein lipase activity, lower intracellular triacylglycerol and glycerol levels, and enhance the release of glycerol into the medium. CLA, especially the t10,c12 isomer, showed its antiobesity effect at 50–200 \( \mu \)M by inhibiting proliferation, suppressing triglyceride accumulation, and inducing apoptosis in 3T3-L1 preadipocyte cultures compared with albumin vehicle or linoleic treatments (98).

Lin et al. (99) found that the t10,c12 CLA isomer inhibited the activity of heparin-releasable lipoprotein lipase (HR-LPL) more strongly compared with the c9,t11 isomer, while both CLA isomers exhibited an inhibitory effect on HR-LPL compared with the LA treatment in 3T3-L1 adipocyte cultures. Yamasaki et al. (100) showed in a rat feeding study that 2% dietary CLA lowered serum leptin level after 1 week compared to the 8% safflower oil supplement control group. Leptin level in perirenal white adipose tissue was also low in animals fed CLA after 12 weeks of
feeding. These findings suggest that the CLA mechanism for body fat reduction in mice, and possibly in other animals, is a result of inhibition of fat transportation and storage in adipocytes coupled with both elevated β oxidation in skeletal muscle and an increase in skeletal muscle mass (71).

D. Bone

Our laboratory recently investigated the effects of CLA isomers on bone modeling in growing male rats (73,74). In our studies with rats, 1% dietary CLA combined with two ratios of n-6/n-3 fatty acids led to differences in CLA enrichment of various organs and tissues. Brain exhibited the lowest concentration of isomers, but bone tissue (periosteum and marrow) contained the highest amounts (73). Both n-3 fatty acids and CLA lowered ex vivo prostaglandin E2 (PGE2) production in bone organ culture. The supplemental CLA isomers also reduced serum insulin-like growth factor type I (IGF-I) concentration and modulated IGF-binding protein (IGFBP) differentially depending on the ratio of n-6/n-3 fatty acids in the diet. Moreover, CLA increased IGFBP in rats given a high dietary level of n-6 fatty acids but decreased IGFBP in rats given a high level of n-3 fatty acids. In tibia, rats given CLA had markedly reduced mineral apposition rate (MAR) (3.69 vs. 2.79 μm/day) and bone formation rate (BFR) (0.96 vs. 0.65 μm/day) in comparison with those not given the CLA supplement (74). Dietary lipid treatments did not affect serum intact osteocalcin or bone mineral content. These results showed that a mixture of CLA isomers at 1% of the diet modulated local factors that regulate bone metabolism and reduced bone formation rates. This response may be due to the total dietary level of CLA or varying effects of individual isomers of CLA on bone biochemistry and physiology.

In a subsequent study with rats, a 0.5% dietary level of CLA was provided with or without beef fat (101). The dietary CLA treatments resulted in total CLA values ranging from 0.27% to 0.43% in the polar lipid fraction and from 2.02% to 3.37% in the neutral lipids in liver, bone marrow, and bone periosteum. We observed that CLA accumulated at a higher concentration in neutral lipids compared with polar lipids consistent with the findings of Ip et al. (56) for rat mammary gland. In rats, the t10,c12-18:2 isomer was incorporated into the phospholipid fraction of tissue lipid extracts at the same extent as was the c9,t11 isomer. The ratio of c9,t11/t10,c12 roughly reflected the isomeric distribution of these CLA isomers in the diet or supplement given to rats. Rat serum osteocalcin, a serum bone formation marker, was decreased in rats given CLA after 12 weeks of dietary treatment. Serum bone-specific alkaline phosphatase activity was also significantly decreased in rats given CLA. The fact that CLA lowered serum bone formation biomarkers, together with our previous finding that CLA lowered ex vivo PGE2 production in bone organ culture and bone formation rate, suggests that some CLA isomers may exert a down-regulatory effect on bone metabolism in growing animals. Future research must be conducted to evaluate the effects of individual CLA isomers on osteoblast function and bone formation.

VI. MECHANISMS OF CLA ACTION

Continued research on CLA has focused on elucidating its mode of action. The CLA isomers have been demonstrated to possess antioxidant properties (54,55,63,64), inhibit carcinogen-DNA adduct formation (58,102,103), induce apoptosis (60), mod-
ulate tissue fatty acid composition and eicosanoid metabolism (61,62,67,68,73,74, 104–106), and affect the expression and action of cytokines and growth factors (68). Since most of the studies mentioned above were conducted with mixtures of several CLA isomers (mainly c9,t11-18:2 and t10,c12-18:2), these proposed mechanisms of action might be specific for individual CLA isomers.

A. Antioxidative Action

Antioxidants originating from both natural and synthetic sources with diverse structures have been known to demonstrate some anticarcinogenic activity (107). Since CLA is proposed to possess antioxidative action, this could explain an important mechanism related to its anticarcinogenic activity. Several studies have been conducted using both in vivo and in vitro systems to clarify the role of CLA as an antioxidant; however, the results obtained from various test systems were conflicting. Ha et al. (54), for the first time, showed that CLA was a potent in vitro antioxidant. In an in vivo study, Ip et al. (55) reported that CLA was as effective as vitamin E in inhibiting the formation of TBARs in the mammary gland. Similar to the results of Ha et al. (54), Ip et al. observed no dose–response relationship in the dietary range of 0.25–1.5% CLA. All doses tested produced a 30–40% inhibition of peroxide formation.

A study with female rats given diets containing different lipids (20% corn oil or lard) or amounts of a fat blend (from 10–20% by weight) with or without 1% CLA (56) showed that CLA reduced malondialdehyde (a peroxidation product) production in rat mammary gland homogenate. In addition, CLA produced a greater reduction in rats fed a diet high in PUFA (corn oil, 35%) than in those given a high saturated fat lard diet (25%). CLA failed to show any inhibitory effect on 8-hydroxydeoxyguanosine (8-OHdG, a marker of oxidative damage to DNA) level in rats. Leung and Liu (108) found that the t10,c12 CLA isomer exhibited a strong antioxidative property compared to the c9,t11 isomer and even \( \alpha \)-tocopherol at a lower concentration of 2 and 20 \( \mu \)M in a total oxyradical scavenging capacity assay. The c9,t11 isomer yielded a weak antioxidant activity at lower concentrations of 2 and 20 \( \mu \)M, but at a higher concentration (200 \( \mu \)M) it performed as a strong pro-oxidant.

Although CLA has been shown to be antioxidative in both in vitro and in vivo studies, the anticancer effect of CLA in these studies cannot be satisfactorily explained based on the current findings. First, the maximal effective concentration of CLA in inhibiting peroxide formation did not agree with the most effective concentration in tumor inhibition (55). Second, neither of the studies demonstrated a dose–response relationship between CLA concentration and its antioxidative efficacy, which is usually true for antioxygenic nutrients studied thus far.

van den Berg et al. (21) reinvestigated the antioxidative property of CLA using a lipid membrane system consisting of 1-palmitoyl-2-linoleoyl phosphatidylcholine. The results of this study indicated that CLA did not show any protective effect under the test conditions and was more susceptible to oxidative damage than LA and comparable to arachidonic acid. In agreement with the findings of van den Berg et al. (21), Banni et al. (109) in a recent study also showed that CLA was more prone to oxidation than LA, and no significant antioxidant effect of CLA was detected in the models tested. It would appear that CLA and its metabolites seem to behave like other PUFAs under conditions of oxidative stress.
In contrast to the antioxidative properties of CLA, results from experiments using human cancer cell lines suggested that CLA, because of its susceptibility to oxidative damage, could behave as a prooxidant in cell culture systems. Moreover, CLA may create an oxidatively stressed environment that is cytotoxic to cultured cells (77,110). O’Shea et al. (110) used CLA dissolved in ethanol, which was added to a human cell culture (MCF-7 breast cancer line and SW-480 colon cancer cell line) media at concentrations of 0, 5, 10, 15, 20, and 30 ppm and incubated for 4, 8, and 12 days. The CLA treatment at 20 ppm increased lipid peroxidation and induced the expression and activity of antioxidant enzymes (superoxide dismutase, catalase, and glutathione peroxidase) in both cell lines. At 20 ppm, CLA also reduced \(^3\)H-leucine incorporation into protein by 83–91% and \(^3\)H-uridine and \(^3\)H-thymidine incorporation into RNA and DNA by 49–91% and 86–98%, respectively, compared with untreated control cells. O’Shea et al. (111) showed that milk fat enriched with CLA (primarily the triglyceride-bound c9,t11 isomer) induced the activities of superoxide dismutase (SOD), catalase, and glutathione peroxidase (GPx) in MCF-7 human breast cancer cell cultures and inhibited the growth and proliferation of these cells. Apparently, at least in part, the induced antioxidant enzyme system failed to protect these cells from peroxidative cytotoxicity. The controversial effect of CLA as an antioxidant or prooxidant agrees with previous knowledge that the balance between antioxidant and prooxidant activity is known to be a complex function dependent on the concentration of the testing material and oxygen partial pressure (112).

Basu et al. (113,114) reported the investigations of the urinary levels of 8-iso-PGF\(_{2\alpha}\), a major isoprostane, and 15-keto-dihydro-PGF\(_{2\alpha}\), a major metabolite of PGF\(_{2\alpha}\), as indicators of nonenzymatic and enzymatic lipid peroxidation after dietary supplementation of CLA in healthy and obese human subjects. A significant increase of both 8-iso-PGF\(_{2\alpha}\) and 15-ketodihydro-PGF\(_{2\alpha}\) in urine was observed after 3 months and 1 month of daily CLA intake (4.2 g/day).

B. Biochemical and Physiological Actions

Another potential action of CLA in preventing carcinogenesis is on DNA adduct formation. For example, dietary CLA (1%, 0.5%, 0.1% diet) inhibited PhIP (a mammary carcinogen)—DNA adduct formation in F344 rat liver and white blood cells in a dose-dependent manner (102). Similar findings were also reported by Liew et al. (58) that when F344 rats were given CLA (0.5% diet equivalent) by gavage, the treatment significantly reduced IQ (2-amino-3-methylimidazo[4,5-f]quinoline)—DNA adduct formation in the colon. Schut et al. (103) used the same rat model to study CLA effects on both PhIP and IQ and found that CLA (0.1–1%) inhibited PhIP-DNA adduct formation in mammary gland and the colon. It was concluded that CLA could be a potential chemopreventive agent against PhIP- or IQ-induced tumors in rodents.

The isomers of CLA might also initiate apoptosis to protect mammary gland cells from chemically induced carcinogenesis. Ip et al. (60) showed in a primary culture of rat normal mammary epithelial organoids (MEOs) that CLA (0–128 \(\mu\)M), but not LA, inhibited growth of MEO which was further shown to be mediated by a reduction in DNA synthesis and a stimulation of apoptosis. Ip et al. (115) showed that CLA was able to increase chromatin condensation and to induce DNA laddering.
—both evidence of apoptosis—in a rat mammary tumor cell line, indicating a potential mechanism of action of CLA. By inducing apoptosis in mammary gland epithelial cells, CLA could prevent breast cancer by reducing mammary epithelial cell density and inhibiting the outgrowth of initiated MEO.

Park et al. (116) showed that LA stimulated MCF-7 breast cancer cell growth, whereas CLA was inhibitory. The LA stimulated phospholipase C (PLC) activity and tended to increase membrane protein kinase C (PKC) activity. However, CLA supplementation did not modify membrane PLC or PKC activity. PGE₂ production was not influenced by LA or CLA addition in this experiment.

Isomers of CLA may exert their biochemical and physiological effects by modulating tissue or cellular eicosanoid metabolism (61,62,67,73,74,104–106). In studies to determine the physiological action of CLA, investigators found reduced PGE₂ [a cyclooxygenase (COX)-catalyzed product of arachidonic acid] concentration in rat serum and spleen (67,104). The amount of PGE₂ in cultured keratinocytes (106,117) and in ex vivo bone organ culture (73) were lowered with CLA treatment in cells and rats, respectively. Igarashi and Miyazawa (118) studied the inhibitory effect of CLA on the growth of a human hepatoma cell line, HepG2, and found that CLA's effect was interfered significantly by addition of LA or arachidonic acid. Addition of antioxidants, such α- tocopherol and BHT, did not diminish CLA's inhibitory effect on cell proliferation, indicating that CLA's effect was not mediated by induction of lipid peroxidation but rather by changes in fatty acid metabolism. Moreover, CLA reduced the level of leukotriene B₄ (LTB₄) from the exudates of cells (67) and thromboxane A₂ (TXA₂) (another cyclooxygenase-catalyzed product of arachidonic acid) in platelet suspension (119). Indirectly, CLA was found to suppress the growth of human MCF-7 breast cancer cells in culture synergistically with NDGA (a lipoxygenase inhibitor) (61), suggesting an inhibitory effect of CLA on the lipoxygenase pathway.

The mixed isomers of CLA were detected in numerous tissues examined in animals given a dietary supplement of CLA (73,104). In one of our experiments, four groups of male rats were given a basal semipurified diet (AIN-93-G) containing 70 g/kg of added fat for 42 days (73,74). The fat treatments were formulated to contain two levels of CLA (0% and 1% diet) and either n-6 (soybean oil having a ratio of n-6:n-3 fatty acids of 7.3) or n-3 fatty acids (menhaden oil + safflower oil having a ratio of n-6 to n-3 fatty acids of 1.8) following a 2 × 2 factorial design. The CLA isomers analyzed by GC were found in all rat tissues analyzed although their concentrations varied with brain exhibiting the lowest concentration of CLA isomers but bone tissues (periosteum and marrow) containing the highest amounts. Dietary CLA decreased the concentrations of 16:1n-7, 18:1, total monounsaturates, and n-6 fatty acids, but increased the concentrations of n-3 fatty acids (22:5n-3 and 22:6n-3), total n-3, and saturates in the tissues analyzed. Ex vivo PGE₂ production in bone organ culture was decreased by the n-3 fatty acid and CLA treatments. In a subsequent study with rats, feeding 0.5% CLA resulted in total CLA concentrations ranging from 0.27% to 0.43% in the polar lipid fraction and from 2.02% to 3.37% in the neutral lipids in liver, bone marrow, and bone periosteum.

When CLA is incorporated into membrane phospholipids (induced by feeding a dietary supplement), it may compete with arachidonic acid and is likely to inhibit eicosanoid biosynthesis (74,120,121). The reduction in the amount of n-6 fatty acids in peritoneal exudate cells and splenic lymphocyte total lipids by CLA seemed to be
responsible, at least in part, for the reduced eicosanoid levels (67). CLA could also affect the lipoxygenase pathway to reduce product formation. Cunningham et al. (61) showed that the addition of CLA and NDGA to MCF-7 cells resulted in synergistic growth suppression, suggesting that CLA effects were mediated through lipoxygenase inhibition.

The effect of dietary CLA on eicosanoid metabolism could be twofold. First, CLA isomers could directly compete with substrate concentration and activity or expression of cyclooxygenase or lipoxygenase. Second, CLA could be further desaturated and elongated to its 20-carbon equivalent and exert its effect by competing with other 20-carbon fatty acids, mainly arachidonic acid and eicosapentaenoic acid, to reduce the production of their corresponding eicosanoids. Experiments to evaluate the effects of CLA isomers on eicosanoid biosynthesis have not yet been conducted. We hypothesized that CLA depressed arachidonate-derived eicosanoid biosynthesis since dietary sources consistently reduced ex vivo PGE₂ production in rat bone organ culture and liver homogenate (74). The reduction in PGE₂ by CLA might be explained as a competitive inhibition of n-6 PUFA formation that results in lowered substrate availability of cyclooxygenase. Although there was a trend of reduced arachidonic acid concentration in bone tissues, the dramatic decrease in ex vivo PGE₂ production in bone organ culture could not be satisfactorily explained by a lack of substrate (74).

The biosynthesis of PGE₂ in bone (cells of the osteoblast lineage) is highly regulated by local and systemic factors (122–124). Fatty acids have been shown to modulate the expression and activity of this key enzyme. For example, Nanji et al. (125) showed that saturated fat reduced peroxidation and decreased the levels of COX-2, the inducible form of COX, in rat liver. In a rat dietary study on colon tumorigenesis, a high-fat corn oil diet (rich in n-6 fatty acids) up-regulated COX-2 expression, but a high-fat fish oil diet (rich in n-3 fatty acids) inhibited it; however, expression of COX-1, the constitutive enzyme, was not affected (126). We speculate that CLA may influence PGE₂ production through the COX enzyme system, more likely on COX-2, to exert its physiological effects in bone and other tissues to influence bone metabolism as well as cancer.

Sebedio et al. (105) reported that CLA may be further desaturated and elongated to form conjugated 20:4 isomers that might block the access of arachidonic acid to COX. The unusual 20:4 isomers derived from CLA might also affect the activity of the COX enzymes. Further study with CLA is needed to confirm if its isomeric analogs alter COX activity and expression as a primary mechanism of action and potential role in controlling cancer and inflammatory disease.

The isomers of CLA were shown to modulate the expression and activity of cytokines and growth factors. Buison et al. (127) reported that 1% CLA in a 40% (wt/wt) fat diet lowered circulating IGF-I level in obese female Wistar rats. We reported that dietary CLA lowered basal and lipopolysaccharide (LPS) stimulated interleukin-6 production and basal tumor necrosis factor production by resident peritoneal macrophages in rats (68). Furthermore, CLA reduced the release of LTB₄ (67), a strong bone resorption factor (128), from peritoneal exudate cells and splenic cells in response to the dietary CLA levels. Assuming that CLA would have similar effects on these cytokines in bone, together with the fact that CLA reduced the production of PGE₂ in bone tissue, one could hypothesize that a proper dietary level, the anti-
inflammatory effects of CLA would be beneficial for the treatment of inflammatory bone diseases.

In our laboratory, the dietary CLA effects on serum concentrations of IGF-I and IGFBPs, and their subsequent impact on bone modeling, were examined in male rats (74). The level of IGFBP in serum of rats was altered by n-6 and n-3 fatty acids, but CLA had variable effects. Interestingly, CLA increased IGFBP level in rats given a high dietary level of n-6 fatty acids but reduced it in those given a high level of n-3 fatty acids. Rats given the n-3 fatty acids had the highest serum level of IGFBP-3. This study also showed that CLA decreased the amount of IGF-I mRNA in liver of rats given n-3 fatty acids. In liver of rats, the expression of IGF-I mRNA appeared to be up-regulated by n-3 fatty acids and down-regulated by CLA. The lowering effect of CLA on growth factors was associated with reduced mineral apposition rate and bone formation rate in the tibia. These results showed that dietary PUFA type (and level) and CLA modulate growth factors that regulate bone metabolism and other aspects of health.

Different CLA isomers may also have their own unique mechanisms of action. Although limited information is available on the biological activity of individual CLA isomers, the data suggest that individual isomers exert different biological effects. Currently, the c9,t11 isomer has been shown to be effective in reducing mammary carcinogenesis and the t10,c12 isomer is more potent in inducing body compositional change. de Deckere et al. (129) treated hamsters with CLA preparations containing relatively pure c9,t11, t10,c12, or a mixture of both isomers and showed that the t10,c12 isomer was the most active form of CLA in inducing biological effects, such as increasing liver weight, decreasing fat deposition, and lowering LDL cholesterol. The authors concluded that t10,c12 appeared to be the physiologically active CLA isomer and the natural c9,t11 had little or no effect on lipid metabolism in hamsters.

Studies performed on mouse and cell culture also indicate differences between the two major CLA isomers in their clearance rate from the body. In skeletal muscle of mice treated with dietary CLA supplements, the t10,c12 isomer was cleared significantly faster than the c9,t12 isomer (94). Yotsumoto et al. (130) recently showed that t10,c12, but not the c9,c11 isomer or LA inhibited cellular triacylglycerol synthesis and reduced apolipoprotein B secretion in HepG2 cell cultures. In the same study, the c9,c11 isomer inhibited cholesteryl ester synthesis but to a lesser extent than the t10,c12 isomer (130). Information on the effects of individual CLA isomers is inadequate at present because of the limited supply of pure CLA isomers. With the advance in techniques for CLA preparation, individual CLA isomers will be available for future research.

C. Immune Function Modulation

Though CLA appears to have an anti-inflammatory effect, Turnock et al. (131) showed that feeding mice CLA for up to 4 weeks does not compromise their immune function against *Listeria monocytogenes*, an intracellular pathogen.

In a study evaluating the effect of CLA on immune competence, early-weaned pigs were given 0–2% of a dietary CLA supplement (132). On day 42, CLA induced a linear increase in percentages of CD8+ cytotoxic/suppressor T cells, indicating that CLA could be effective in inhibiting disease-associated growth suppression in pigs.
Yamasaki et al. studied CLA and antibody production in vivo and in vitro (133). In CLA-fed (0.05–0.5%) Sprague–Dawley rats, the production of IgG, IgM, and IgA in spleen lymphocytes was dose-dependently increased, whereas the serum concentration of these immunoglobulins were not affected. In an in vitro assay with spleen lymphocytes, CLA at 100 μM suppressed Ig production (133).

Dietary CLA supplement of 0.25 g per 100 g of diet for a week significantly reduced histamine and PGE₂ release from female Hartley guinea pig trachea tissue superfusate when sensitized with antigens, indicating a reduced release of some inflammatory mediators during type 1 hypersensitivity reactions (134). Feeding healthy young women subjects 3.9 g/day CLA did not change the indices of immune status, such as number of circulating white blood cells, granulocytes, monocytes, lymphocytes, and their subsets; lymphocyte proliferation in response to phytohemagglutinin; and influenza vaccine, serum influenza antibody titers, and DTH response (20,51).

VII. POTENTIAL ADVERSE EFFECTS OF CLA

Since CLA has become a widely advertised nutritional supplement for human use (135), it is important to study both its positive and potential negative effects on health using cell culture or animal models. Our recent dietary studies using an isomeric mixture of CLA revealed a negative effect on rat bone metabolism (74,101). Rats given 1% CLA in the diet demonstrated decreased mineral apposition rate and bone formation rate in the tibia compared with rats not given CLA. In a follow-up experiment with rats, a lower level of CLA (0.5% diet) reduced serum osteocalcin and bone-specific alkaline phosphatase activity, both biomarkers of bone formation, after 12 weeks of dietary treatment (136). The negative effect of CLA was likely due to a high dietary level of isomeric mixtures that do not reflect the usual food sources of CLA.

In a recent study by Belury et al. (137), CLA displayed the typical peroxisome proliferation response in rodent liver. Peroxisome proliferators may enhance tumorigenesis in liver, testes, and pancreas by acting as promoters, resulting in enhanced cell proliferation, altered cell differentiation, and inhibition of apoptosis in initiated cells (137). This response might suggest that the chemoprotective effect of CLA in extraplastic tissues (55,56,58,76) may be at the expense of enhanced hepatocarcinogenesis. Jones et al. (138) also reported that CLA lowered blood LDL cholesterol but increased VLDL cholesterol and resulted in liver hypertrophy in mice.

The results of Belury et al. (137) indicate that the peroxisome proliferation response to CLA may be greater in mice than in rats. These data suggest a species difference in the response to CLA, and such information might be relevant to a proper risk assessment for human consumption of isomeric mixtures of CLA. Thus far, the information on CLA action in humans is still very limited; additional research is necessary to clarify safety issues related not only to isomeric supplements of CLA but also to individual isomers that could behave differently from each other in various biological systems and physiological conditions.

Many CLA isomers present in commercial CLA supplements for human use do not exist in natural food products. Furthermore, even for the naturally occurring CLA isomers, human consumption without dietary supplementation is normally at a very low level. The possible negative effects of CLA at trace levels could not be easily detected. Therefore, before promoting the human use of dietary CLA supple-
ments, thorough examination of the effects of individual CLA isomers is necessary to protect consumers from potential detrimental effects. Together with the fact that CLA lowered serum bone formation biomarkers and bone formation rate, we believe that specific research must be implemented to evaluate the safety of these unique fatty acids.

REFERENCES


Dietary Fats and Obesity

DOROTHY B. HAUSMAN, DANA R. HIGBEE, and BARBARA MULLEN GROSSMAN

The University of Georgia, Athens, Georgia

I. INTRODUCTION

Obesity is variably defined as having a body weight of more than 20% over ideal body weight or a body mass index [weight (kg)/height (m)²] of 30 or more (1). Obesity is associated with many chronic diseases and alterations in physiological function, including cardiovascular disease, hypertension, diabetes mellitus, gallbladder disease, and certain types of cancer (2). Obesity is a major public health problem in the United States and Europe and is becoming increasingly important in many other areas of the world (3). The prevalence of obesity in adults in the United States has increased by 30% or more in the past decade, with increases in both genders and in all ethnic and racial populations and age groups (4). It is now estimated that in the United States more than one-third of the total adult population and one-half of the adult African American and Hispanic population are obese (5).

The etiology of human obesity is quite complex, involving genetic, metabolic, behavioral, and environmental factors. Although obesity is believed to have a strong genetic component (6), the increased incidence of obesity in specific population groups undergoing westernization indicates the importance of dietary and lifestyle changes in the manifestation of this disease (7,8). Among dietary factors, both total energy intake and fat intake are significantly correlated with body mass index in these population groups (7). However, increased intake of fat energy is associated with a greater per unit increase in body mass than is increased intake of energy from nonfat sources. Therefore, much attention regarding dietary influences on obesity development or prevention has focused on high-fat diets.
II. DIETARY FAT AND BODY WEIGHT

An effect of dietary fat on body weight has been well documented, especially in experimental animals. Warwick and Schiffman (9), in a thorough examination of the literature, reviewed the effects of diet type (fat vs. carbohydrate) and caloric density on caloric intake and weight gain reported in animal studies. They suggested that the caloric density of the diet plays a major role in promoting weight gain. If the density is 4.5 kcal/g or more, excess weight gain is more likely to occur. In several instances, an increase in body weight occurs without an increase in food intake.

Human studies, because they are more difficult to control, are less prevalent than studies with animals. However, human studies also demonstrate that dietary fat promotes weight gain. Lissner et al. (10) demonstrated that subjects consuming a high-fat diet gained weight whereas those eating a low-fat diet lost weight. Others (11,12) have reported that switching from a high-fat to a low-fat diet promotes relative weight loss. Finally, indirect evidence provided from epidemiologic studies shows positive correlations between body weight and fat intake (13).

III. DIETARY FAT AND FOOD INTAKE

A. Does Dietary Fat Affect Food Intake?

Dietary fat can have a profound effect on energy balance and, ultimately, body weight. Studies designed to investigate the effect of dietary fat on food intake generally employ one of two approaches: short-term studies examining the influence of dietary fat on meal size or frequency, and longer term investigations of the effects of dietary fat on energy intake over days or weeks. Short-term effects are typically examined in one of two ways: (a) fat is given as a “preload” and subsequent feelings of hunger or food intake is reported; or (b) fat is given as part of a mixed diet and concurrent food intake is measured. In the first instance, “satiety,” i.e., the ability of a substance to suppress further eating, is measured by the time elapsed, or amount of food eaten at the next meal. In the second case, “satiation” is assessed and is defined as the size of the current meal (14).

Many studies have been carried out examining the effect of a “preload,” often in the form of a liquid, on subsequent intake, typically over a short time period. Others have investigated the effect of high- or low-fat meals or snacks, or meals supplemented with fat or carbohydrate, on subsequent feelings of fullness and food intake. The results of these studies have been quite variable [for review, see Blundell et al. (15)]. Many reports indicate that with a preload, individuals do not fully compensate for the calories ingested, i.e., they do not reduce their intake in accurate proportion to the calories previously consumed (16–19). This seems to be truer for fat versus carbohydrate preloads (18–24), though some studies have indicated that fat has satiety value equal to carbohydrate (25,26) or even greater than carbohydrate (27,28). In addition, compensation seems to occur initially but then decreases over time (25–27).

It is important that the volume, sensory characteristics, and protein content of the preloads be similar when investigating the satiety effect of fat. A number of studies have not controlled for all of these factors (17,24,29,30). However, when preloads are similar, investigators have found that individuals vary substantially in their response to preloads and that body weight may play a role in these responses.
For example, some have reported that males who are of normal weight and not concerned about their body weight or food intake ("unrestrained eaters") appear to compensate adequately for the caloric content of a preload (24). Porrini et al. (31) report that a high-protein food given as a snack 2 hours before a meal exerts a higher effect on both intrameal satiation and postingestive satiety than a high-fat snack. When a first course is consumed as part of a meal, the sensory characteristics of the foods play an important role in controlling subsequent food intake. Others (20) have shown that in obese individuals, fat exerts only a weak action on satiety. Data such as these have led to the speculation that obesity may be the result of an insensitivity to satiety signals generated by ingestion of fat.

In studies of satiation, dietary fat is an integral variable of the test diet(s). Studies show that caloric intake is greater with dietary fat than with carbohydrate. When subjects are fed diets in which fat, fiber, and simple sugars are manipulated to obtain low energy versus high energy diets, energy intake is greater on the high energy as compared with the low energy diet (32). Caputo and Mattes (33) reported that individuals consuming high-fat meals consume more calories than those consuming high- or low-carbohydrate or low-fat meals. Others (34) have shown that individuals consuming high-fat or high-carbohydrate diets for 1 week consume more calories on the high-fat versus high-carbohydrate diets. This hyperphagic effect of dietary fat has been observed in many studies (35–39).

The excessive intake of dietary fat primarily occurs during a meal to increase meal size rather than between meals to increase meal frequency (37). Overconsumption is particularly high when fat is combined with alcohol (40). Over the long term, many investigators have shown that this passive overconsumption of dietary fat can lead to obesity [for review, see (37,41)].

Relatively few long-term studies on humans have been conducted because of difficulties such as inability of the investigator to control for the subjects' current or past food intake, activity level, or genetic background. However, Kendall et al. (11) reported that women consuming either a low- or high-fat diet for two separate 11-week periods consume more calories on the high-fat diet. Lissner et al. (10) also showed that subjects eating either a low-, medium-, or high-fat diet for 2 weeks consume the most calories on the high-fat diet.

In addition, animal studies have shown that the type of fat may influence the satiation effects of dietary fat. In both chickens (42) and rats (43), medium chain triglycerides have been shown to have a greater satiating effect than long chain triglycerides. Rolls (44) and Stubbs (45) both report that in humans, substitution of long chain triglycerides with medium chain triglycerides depresses food intake. Also, the physical form as well as the type of fat contributes to its satiation effects (27,46).

Overeating occurs with diets containing saturated fats (47–50) as well as mixed fats (51–54). Recently, Lawton et al. (55) showed that in human subjects polyunsaturated fatty acids may exert a stronger control over appetite than monounsaturated or saturated fatty acids.

Some have suggested that the hyperphagic effect of high-fat diets is not due solely to the fat but is influenced by the presence of carbohydrate and overall caloric density (56). Ramirez et al. (56) fed rats diets varying in carbohydrate, fat, cellulose, or caloric density and found that energy intake varies directly as a function of caloric density regardless of the fat or cellulose content of the diets. They concluded that high levels of fat, carbohydrate, and energy interact to produce overeating in animals.
fed high-fat diets. In support of this hypothesis, Emmet and Heaton (57) examined food records from 160 subjects who had weighed their food for 4 days. They reported that an increase in refined sugar intake is associated with a linear increase in the intake of fat combined with carbohydrate. This suggests that refined sugar may act as a vehicle for fat intake by increasing fat palatability.

Some have reported that there is an inverse relationship between consumption of sugar and fat, terming it the “see-saw” effect. When consumption of fat and sugar are expressed as a percentage of total intake, then this inverse relationship is present. However, when expressed in absolute terms, there is a positive relationship between dietary fat and carbohydrate intake (58).

B. Mechanisms for Fat-Induced Food Intake

1. Caloric Density

The higher caloric density of many high-fat diets may play a role in inducing this hyperphagic response (36,37,59). In humans, Duncan et al. (32) reported that adult subjects eat almost twice as many calories on a high density diet compared to a low density diet. In studies using experimental animals, hyperphagia is typically observed only when the caloric density of the diets is high, greater than approximately 5.8 kcal per gram of diet [for review, see Warwick and Schiffman (9)]. In addition, when caloric density is constant, rats fed a diet high in corn oil have a caloric intake similar to that of animals fed low-fat diets (56,60).

Several investigators have reported that a preference develops for a flavor that is paired with a high number of calories versus one paired with a low number of calories. A study by Johnson et al. (61) indicated that children report increased flavor pleasantness due to association with a high density of fat calories. In rats, a flavor associated with corn oil consumption is preferred over a flavor that is not paired with oil (62,63). In studies in which oils or high-fat foods are given, animals often initially do not consume greater quantities of the fat. However, over time the rats do consume more of the fat as they learn about the associated postingestive consequences (i.e., greater caloric value) (64–68). In contrast, others have suggested that that dietary fat is overconsumed even when compared with an isoenergetic carbohydrate diet of similar palatability (69,70). Further, Lucas et al. (71) report that, relative to an isocaloric high-carbohydrate diet, the postingestive effect of high-fat diets stimulates overeating and conditions a stronger flavor preference in rats, suggesting that some quality in fat per se may be inducing intake.

2. Stomach Distention

Differences in stomach distention due to dietary fat versus carbohydrate or protein may also account for its hyperphagic effect (9). Warwick and Schiffman (9) suggested that because of the greater caloric density of dietary fat, a high-fat meal has a smaller volume than an isocaloric high-carbohydrate meal, resulting in less stomach distention. This lesser distention would lead to an attenuation of satiety signals. In addition, Cunningham et al. (72) reported that the rate of stomach emptying increases when a high-fat diet is habitually consumed.

Dietary fat has also been found to influence relative consumption of carbohydrate and protein. Crane and Greenwood (73) allowed rats to select from either high-carbohydrate or high-protein diets. Half of the diets contained 20% soybean oil as
the fat component and the other half contained 20% lard as the fat component. Animals selecting from the soybean oil diets consumed more carbohydrate and less protein than animals choosing from the lard-based diets. Grossman et al. (74) further showed that rats gavaged with either beef tallow or corn oil 2 hours before selecting from either high-carbohydrate or high-protein diets consume less carbohydrate and more protein if given tallow versus corn oil. They (74) also demonstrated that the hepatic vagus must be intact for this selection to occur and that mercaptoacetate (fatty acid oxidation inhibitor) can blunt the effect.

3. Metabolic Signals

In an attempt to understand the mechanisms underlying the effects of dietary fat on appetite, investigators have examined absorptive or postabsorptive responses to dietary fat. In particular, fatty acid oxidation has been studied as a possible satiety signal that is generated in response to food (i.e., fat) consumption. Scharrer and Langhans (75) demonstrated that in rats, consumption of a high-fat diet can be stimulated by inhibiting fatty acid oxidation with 2-mercaptoproacetate. Friedman et al. (76) also reported a stimulation of food intake when fatty acid oxidation is inhibited with methyl palmoxyrate. This feeding effect, labeled “lipoprivic feeding,” has been shown to be impaired by hepatic vagotomy (77) and subdiaphragmatic vagotomy (78). In addition, Ritter and Taylor (79) reported that capsaiacin can block this effect, implying that vagal sensory neurons appear to be involved in lipoprivic feeding. Studies utilizing brain lesions indicate that lipoprivic feeding involves the lateral parabrachial nucleus and possibly the area postrema/nucleus of the solitary tract (80). Type of fat may influence this response as Wang et al. (81) reported that food intake is stimulated by mercaptoacetate in rats given corn oil but not tallow diets.

Other investigators have examined aspects of fat metabolism as potential satiety signals by administering fat metabolites centrally or peripherally. Arase et al. (82) reported that intracerebroventricular infusions of \( \beta \)-hydroxybutyrate (\( \beta \)-OHB) reduce food intake in Sprague–Dawley or Osborne–Mendel rats consuming either a high- or a low-fat diet. However, such infusions do not reduce food intake in S5B/PI rats, rats that are resistant to weight gain when consuming high-fat diets. Peripheral injections of \( \beta \)-OHB decrease food intake in S5B/PI rats by not Osborne–Mendel animals, and glycerol has no effect in either strain (83). Peripheral injections of glycerol in Wistar rats (84) and \( \beta \)-OHB in Sprague–Dawley rats (85) also decrease food intake.

Fat, when infused into the intestine, can suppress hunger, induce satiety, or delay gastric emptying (16,86–88). However, as stated previously, dietary fat appears to be overconsumed and often can lead to obesity. The disparate effects of intraintestinal infusions of fat versus dietary fat have been termed the “fat paradox” (89). It has been suggested that high-fat foods have a very high palatability and orosensory stimulation (90–92), leading to overconsumption before the nutrients can even enter the intestine to generate satiety signals. In support of this hypothesis, satiety signals potentially arising from postabsorptive metabolic events appear to be blunted from fat in comparison with other nutrients. For example, it has been reported that carbohydrate and protein consumption is followed by an increase in their oxidation (93–95), whereas oxidation of fat is not generally stimulated until 3–7 days following consumption of a high-fat diet (96,97). Furthermore, it has been reported that fat oxidation is especially limited in obese as compared with lean individuals (34,98).
4. Effects of Hormones and Pharmacological Agents

Use of hormones and pharmacological agents has further illuminated factors that may play a role in the effect of dietary fat on food intake. Pancreatic procolipase is a cofactor for lipase, an enzyme necessary for proper fat digestion. A pentapeptide produced by the cleavage of procolipase, Val-Pro-Asp-Pro-Arg, or enterostatin, has been shown to reduce food intake in rats (99), especially when consuming a high-fat diet (100). Peripheral injection or intracarotid injection (101), or injection into the lateral ventricle (102), suppresses fat intake in fat-adapted rats, suggesting both a gastrointestinal site and a central site of action. Also, high-fat feeding and cholecystokinin-8 (CCK-8) increase intestinal enterostatin levels (103). Lin et al. (104) report that β-casomorphin 1–7 stimulates intake of a high-fat diet in rats and this effect is inhibited by enterostatin or naloxone.

Much research has been focused on CCK and its effects on satiety [for review, see Smith and Gibbs (105)]. It has been well documented that consumption of fat stimulates the release of CCK, activating receptors in the stomach. This signal is transmitted along the vagus to the nucleus of the solitary tract, where it is forwarded to the hypothalamus. Vagotomy can block these effects of systematically administered CCK (106), as can CCK-A receptor antagonists (107).

Morphine has also been shown to have specific effects on intake of dietary fat. Rats given morphine injections subsequently increase fat intake while suppressing carbohydrate intake when given separate sources of macronutrients (108–110) or mixed diets (111). Continuous infusion of morphine also stimulates fat intake (112). An opioid agonist, butorphanol, also increases consumption of a high-fat diet (54). Administration of opioid antagonists suppresses fat intake with little effects on protein or carbohydrate intake (113,114). In humans, opioids also appear to play a role in regulating fat intake. Opiate antagonists cause a decrease in intake of fat calories with little effect on carbohydrate consumption (115–117).

Corticosterone has also been implicated in the regulation of fat intake. Castonguay et al. (118,119) reported that adrenalectomy reduces total caloric intake in rats, particularly fat intake, and that corticosterone can restore the fat consumption. Others (120) have reported that the type 1 adrenocorticoid receptor mediates corticosterone’s effect on fat appetite. However, others (121) have suggested that corticosterone acts to enhance carbohydrate rather than fat intake. It has been suggested (122) that the differing levels of micronutrients added to the diets in these studies may account for these disparate findings.

Investigators have also reported that the peptide galanin influences appetite for fat. Leibowitz (123) first demonstrated that galanin stimulates fat intake, especially at the end of the nocturnal cycle. Galanin is thought to work in concert with norepinephrine, which is colocalized with galanin in paraventricular neurons (124). Smith et al. (125) report that centrally injected galanin induces fat intake only in fat-preferring rats, i.e., baseline feeding preferences are important in determining the feeding response to galanin.

Leptin, the gene product of ob gene, is shown to regulate body fat in mice and is produced in human adipose tissue as well [for review, see Harris (126)]. Reports as to its relationship with fat intake in humans indicate that it is negatively associated with fat intake. Havel et al. (127) report that in women, high-fat/low-carbohydrate meals result in a lowering of 24-hour circulating leptin concentration. Other research
shows that there is a negative correlation between leptin levels and dietary fat (7-day records), when controlling for body weight (128). Finally, Niskanen et al. (129) indicate that serum leptin concentrations in obese humans are inversely related to dietary fat intake. In mice, however, high-fat feeding or a high-fat diet increases serum leptin levels (130,131).

IV. DIETARY FAT AND METABOLISM

Obesity is the final result of increased deposition of fat through increased de novo lipogenesis and/or increased fatty acid esterification relative to lipolysis and oxidation. In the next sections, the critical literature that characterizes the role of dietary lipid in altering these metabolic events will be reviewed.

A. Influence of Dietary Fat on Lipogenesis

Dietary lipid level influences the rate of lipogenesis. Early studies showed that de novo synthesis of fatty acids is decreased by high dietary lipid level (132,133). Two key enzymes in the lipogenic pathway, fatty acid synthetase and acetyl CoA carboxylase, are reduced in animals receiving a high-fat diet. In addition, the pentose phosphate pathway and malic enzyme, both of which provide reducing equivalents for de novo lipogenesis, are also influenced by dietary lipid level. Malic enzyme, the pentose phosphate pathway, and the rate-limiting enzyme in this pathway, glucose-6-phosphate dehydrogenase, are decreased in rats fed a diet containing high levels of dietary fat and increased in diets high in carbohydrate (134–137).

The lipid composition of the diet can cause a shift in the source of stored lipid. On a diet rich in lipids, rat adipose tissue fatty acids come mainly from dietary fat, whereas on a high-carbohydrate diet the fatty acids come from hepatic lipogenesis (138). In genetically obese rats, both hepatic and adipose cell lipogenic rates are decreased by a high-fat diet. Nonetheless, the animals still deposit more fat because of the increased uptake of fatty acids from the diet. Some studies in human subjects suggest a minor role of hepatic lipogenesis in energy balance but do not address the issue of extrahepatic lipogenesis [for review, see Hellerstein et al. (139)]. Using a different technique, other investigators suggest that adipose tissue may account for up to 40% of whole-body lipogenesis when comparing subjects fed a high-carbohydrate diet to subjects fed a high-fat diet after a period of energy restriction (140).

The inhibition of lipogenesis is not uniformly influenced by different fatty acids. For example, unsaturated fatty acids are better at inhibiting de novo lipogenesis than are saturated fatty acids (141–143). The inhibitory effects of unsaturated fatty acids are further influenced by fatty acid chain length, degree of unsaturation, and double-bond location. To inhibit genes that code for fatty acid synthesis, the fatty acid must contain a minimum of 18 carbons and possess at least 2 conjugated double bonds (144). In addition, greater inhibition of lipogenesis is observed with n-3 as compared with n-6 polyunsaturated fatty acids (PUFAs) (137,145–147). PUFA-mediated inhibition of lipogenesis may also be influenced by metabolic status as obese mice appear to be somewhat resistant to PUFA feedback control of gene expression (148).

Ingestion of n-6 or n-3 polyunsaturated fatty acids causes a rapid inhibition of the expression/activation of many enzymes involved in lipogenesis [see reviews by
Jump and Clarke (136) and Clarke (137) and a coordinate induction of genes encoding proteins involved in lipid oxidation and thermogenesis (discussed in Sect. IV.C below). PUFA regulation of lipogenic genes is mediated at both the transcriptional level, as for pyruvate kinase, pyruvate dehydrogenase, acetyl CoA carboxylase, and fatty acid synthase, as well as the posttranscriptional level, as for glucose-6-phosphate dehydrogenase [see reviews by Jump and Clarke (136) and Clarke (137)]. There is emerging evidence that the PUFA-induced suppression of lipogenic enzymes is mediated by changes in the expression and cellular localization of the transcription factor, sterol regulatory element–binding protein 1 (SREBP-1) (147,149,150). Overexpression of this transcription factor in the liver leads to a marked elevation in the mRNAs encoding several lipogenic enzymes and to very high rates of de novo lipogenesis (151,152). Consumption of diets containing n-6 (safflower oil) or n-3 (fish oil) fatty acids leads to a decrease in membrane content of SREBP-1 precursor and nuclear content of SREBP and a concomitant reduction in lipogenic gene expression in the liver (147). All effects are greater for the n-3- than the n-6-containing diets (147). Most studies investigating the effects of n-3 fatty acids on SREBPs are limited to lipogenic gene expression in the liver rather than adipose tissue. Whether analogous regulatory processes govern n-3 fatty acid modulation of SREBPs and lipogenic genes in adipose tissue is unknown. Consistent with the inhibitory action of n-3 fatty acids on expression of genes involved in hepatic lipid metabolism, others have observed similar effects in white adipose tissue (153,154). It is tempting to speculate that n-3 fatty acids may also influence gene expression in adipose tissue via regulation of SREBPs.

B. Influence of Dietary Fat on Lipid Uptake

Lipoprotein lipase (LPL) has been called the “gate keeper” enzyme because it controls the rate of uptake of lipid by adipose cells (155). This enzyme is elevated in association with genetic and diet-induced obesity in animals and humans. The ability of n-3 polyunsaturated fatty acids to lower serum triglycerides is thought by some to be due to an action on lipoprotein lipase. Several groups have reported an increase in endogenous LPL activity and post-heparin LPL activity with n-3 fatty acid supplementation in both healthy and hypertriglyceridemic patients (156,157) and experimental animals (158). In contrast, others observed no effect of n-3 fatty acids on LPL activity in humans (159,160) or rats (161). Raclot et al. (153) reported a reduction in LPL expression in retroperitoneal fat depots, but not in subcutaneous fat depots, in rats fed the n-3 fatty acid docosahexaenoic acid (DHA) alone or in combination with the n-3 fatty acid eicosapentaenoic acid (EPA), suggesting site-specific effects. In addition, LPL activity may not necessarily correspond to mRNA levels, as posttranscriptional events, such as glycosylation and binding of LPL to cell surface heparan sulfate proteoglycans, modulate expression and activity of the enzyme (162). Furthermore, while alterations in LPL activity or expression may function in the lowering of serum lipid levels associated with fish oil (n-3 fatty acid) consumption, it is at odds with the finding of reduced adipose tissue mass in animals fed the same diets. Conversely, the reduced obesity associated with diets high in n-3 polyunsaturated fatty acids may be due to the influence of these fatty acids on reducing hepatic fatty acid synthetase activity (158,163) and stimulating fatty acid oxidation (see below).
C. Influence of Dietary Fat on Fatty Acid Oxidation and Energy Expenditure

It has been known since the early days of calorimetry that diets high in fat lower the respiratory quotient, an indicator of increased fatty acid oxidation (164). Fatty acid oxidative rates are dependent in part on chain length and degree of unsaturation. In normal-weight men, fatty acid oxidation rates are highest for lauric acid, followed by polyunsaturated and monounsaturated fatty acids and least for longer chain saturated fatty acids for which oxidation decreases with increasing chain length (165). Dietary lipid stimulation of fatty acid oxidation is thought to act through the sympathetic nervous system (166) and stimulation of carnitine palmitoyltransferase activity (167), both of which are influenced by the source of dietary fat. For example, safflower oil–fed rats have the highest sympathetic activity when compared with coconut oil– or medium-chain triglyceride–fed rats (166). Similarly, feeding fish oil causes a marked increase in carnitine acyltransferase activity in hepatic mitochondria as compared with corn oil feeding (167). Peroxisomal oxidation is also increased by diets containing fish oils when compared with vegetable oils (145). Studies in rodents have demonstrated that the n-3 fatty acids EPA and DHA both stimulate peroxisomal β oxidation in the liver, whereas EPA increases mitochondrial β oxidation (168). Significantly higher mitochondrial and peroxisomal β-oxidative rates in fish oil–fed rats, as compared with those fed palm or safflower oil (169), are presumably due to the up-regulation of genes encoding proteins for mitochondrial and peroxisomal enzymes [see reviews by Clarke et al. (137,170)]. Preferential oxidation of n-3 fatty acids seemingly limits substrate availability for triglyceride synthesis and delivery to adipose tissue, as the oxidative rate of fatty acids is inversely proportional to storage rates (171).

Energy expenditure can be influenced by dietary lipids. In general, high-fat diets have a lower heat increment than diets high in carbohydrate or protein (164). This may lead to a decrease in dietary energy utilization and an increase in body weight gain when fed a high-fat diet. While evidence exists for high-fat diets causing obesity, not all high-fat diets affect energy metabolism and body weight in the same manner. For example, diets high in essential fatty acids result in a lowering of body weight and increase in thermogenin content in rat brown adipose tissue (172). Increased levels of dietary polyunsaturated fatty acids increase hepatic carnitine palmitoyltransferase activity in Zucker obese rats and reduce liver lipid content (173). Diets containing safflower oil cause an increase in thermogenesis (174) and uncoupling protein content (175) of brown adipose tissue in comparison with diets containing beef tallow.

The n-3 fatty acids, present in fish oils, may also induce the expression of uncoupling proteins. As their name implies, uncoupling proteins dissociate mitochondrial oxidative phosphorylation from energy production, leading to energy loss as heat. Three mitochondrial uncoupling proteins (UCPs) have been identified thus far. UCP-1 is expressed predominantly in brown adipose tissue (176), UCP-2 in white adipose tissue and other tissues (177), and UCP-3 in skeletal muscle (178). n-3 fatty acids have been demonstrated to increase expression of UCPs beyond that of other types of dietary fats in several studies. Hun et al. (179) reported that KK-Ay/TaJcl mice fed high-fat diets supplemented with n-3 fatty acids had increased levels of UCP-2 mRNA in white adipose tissue. Similarly, Takahashi and Ide (161) reported
an increase in UCP-1 mRNA in brown adipose tissue of rats fed high-fat diets containing fish oil as compared with safflower oil. Oudart et al. (180) demonstrated an increase in brown adipose tissue mitochondrial guanosine 5’-diphosphate binding with n-3 fatty acid supplementation, but no associated difference in UCP-1. Baillie et al. (170) reported an increase in skeletal muscle UCP-3 mRNA in fish oil–fed rats, which was inversely correlated with a 25% decrease in body fat mass. It has been postulated that one mechanism by which n-3 fatty acids decrease adipose tissue mass in rats and mice is by increasing thermogenesis. However, dissipation of consumed energy as heat does not explain the lack of variance in body weights in some studies.

D. Influence of Dietary Fat on Adipose Tissue Lipolysis

The quantity of fat stored in adipose tissue is determined by the relative rates of the simultaneously occurring metabolic processes of triglyceride synthesis and lipolysis or triglyceride breakdown. The hydrolysis of triglycerides is catalyzed by hormone-sensitive lipase, an enzyme regulated by a complex cyclic AMP–dependent signal transduction cascade (Fig. 1). A variety of hormones and neurotransmitters (e.g., adrenaline, glucagon, noradrenaline) stimulate various components of the signal transduction cascade and thereby increase lipolysis [see review by Vernon (181)]. Insulin, in contrast, modulates the activity of cyclic AMP, thereby reducing lipolytic activity (181).

1. Level of Dietary Fat

There is considerable evidence that both hormone-stimulated lipolysis and the antilipolytic effects of insulin are influenced by the quantity and/or type of dietary fat. However, the effect of dietary fat on lipolysis varies somewhat according to the species and specific adipose tissue depot being studied. For example, feeding rats high levels of dietary fat leads to a decrease in both catecholamine- (182–190) and glucagon- (182) stimulated lipolysis. The effect of high-fat feeding on adipose tissue lipolysis in the rat is believed to be due to changes in β-adrenoceptor number or to an uncoupling between the hormone receptor and adenylate cyclase, rather than to differences in hormone binding (183,184,190). In contrast, several studies have indicated an increase in cyclic AMP–dependent signal transduction and/or lipolytic response with an increase in dietary fat (187,191–195). More specifically, adenylate cyclase activity (191) and isoproterenol-stimulated lipolysis (192) are increased and phosphodiesterase activity is decreased (193) in pigs fed added fat as opposed to control diets. In addition, the effect of high-fat feeding on adenylate cyclase activity and lipolytic response in pigs varies according to the adipose tissue site (191,193). In rats, high-fat feeding leads to an increase in basal and hormone-stimulated lipolytic activity that is positively correlated with fat cell size but is not associated with sympathetic nervous system activity (195). There are few studies regarding the effect of high-fat feeding on adipose tissue lipolysis in humans. However, in one short-term study (7 days), Kather et al. (196) observed no differences in either sensitivity or response to catecholamines in adipose tissue of subjects eating fat-rich as compared with carbohydrate-rich diets.

Fat feeding has been shown to influence the antilipolytic effects of insulin in adipose tissue from several species; however, the results are somewhat inconsistent.
Smith et al. (185) observed a decreased response to the antilipolytic effects of insulin in adipose tissue from high-fat as compared with high-carbohydrate fed rats. However, subsequent studies by Susini et al. (186) and Tepperman et al. (184) failed to observe an effect of dietary fat on the antilipolytic effects of insulin. In pigs, a decrease in the antilipolytic action of insulin in response to added fat feeding is observed in the subcutaneous fat depot but not for the perirenal site (197). In contrast, a greater sensitivity to the antilipolytic action of insulin is observed in adipocytes of human subjects consuming fat-rich as opposed to energy-restricted diets (196).

2. Type of Dietary Fat

Lipolytic response may be influenced by the type as well as the level of fat included in the diet. The composition of dietary fat selectively influences fatty acid and triacylglycerol deposition in adipose tissue (198). In turn, the composition of the fat tissue influences lipid mobilization and release of fatty acids into the circulation. Lipid mobilization from adipose tissue is not a random event, but instead is influenced by chain length, degree of saturation, and positional isomerization of the fatty acids (176,199–202). The relative mobilization of fatty acids from adipose tissue is correlated positively with unsaturation and negatively with chain length (200). The most easily mobilized fatty acids are those with 16–20 carbon atoms and 4 or 5 double bonds, whereas very long chain unsaturated and monounsaturated fatty acids are less readily mobilized (201). Higher rates of lipolysis are also observed with trans as opposed to cis isomers of octadecenoic acid (199). It has been suggested that the decreased fat accumulation in animals fed trans fatty acids may be associated with direct effects of the trans isomer on fat cell metabolism (199).
Adipose tissue from animals fed diets high in polyunsaturated fatty acids generally exhibit a greater lipolytic response to catecholamines and synthetic β-adrenergic agonists as compared with tissue from animals fed diets high in saturated fat (187,203–208). The decreased responsiveness of fat cells from rats fed saturated fat diets is associated with reductions in adenylate cyclase, cyclic AMP phosphodiesterase, and hormone-sensitive lipase activity (203). β-Adrenergic receptor binding is also lower in fat cell membranes from rats fed saturated fat as compared to polyunsaturated fat diets (204,209). The reduction in β-adrenergic receptor binding associated with the saturated fat diet is due to decreased binding affinity rather than to changes in receptor number (204,209). The reduced binding affinity is in turn correlated with a reduction in membrane fluidity in cells from the rats fed the high saturated fat diets (204).

In contrast to the above studies, several investigators have failed to observe an effect of dietary fat type on lipolytic response (188,192,210,211). Lipinski and Mathias (188) observed that norepinephrine-stimulated lipolysis in rat adipocytes is depressed by an increase in fat calories but is unaffected by the degree of saturation of the fat. Likewise, Mersmann et al. (192,210) reported an increase in the number of β-adrenergic receptors, with no change in receptor affinity, in adipose tissue from pigs fed high levels of saturated fat. However, the observed increase in receptor number is not associated with alterations in β-adrenergic agonist–mediated function (including lipolysis). Recently, Portillo et al. (212) observed that under energy-controlled feeding conditions, various dietary fat regimes caused major changes in adipose tissue phospholipid composition; however, such modifications did not lead to important changes in lipolysis. Portillo et al. (212) suggested that some changes in adipose tissue fatty acid composition may have little effect on overall physiological function.

3. Relationship to Obesity

Although the evidence is not totally consistent, it appears the rate of lipolysis may be influenced to some extent by both the level and the type of dietary fat. A reduction in lipolytic rate as is commonly observed in response to high-fat (particularly saturated fat) feeding could lead to an increased retention of stored triglycerides and thereby contribute to the development of obesity. However, this response may vary considerably according to species, age, adipose tissue depot site, and adipose tissue fatty acid composition. The relative contribution of lipolytic alterations to diet-induced obesity and the specific regulatory components of the lipolytic signal transduction cascade influenced by alterations in the level or type of dietary fat remain to be elucidated.

E. Influence of Dietary Fat on Insulin Action

An association between high-fat diets and impaired insulin action has been observed in numerous in vivo and in vitro studies. Early studies in human subjects suggested that diets high in fat lead to a reduction in glucose tolerance (213,214). More recent epidemiologic data suggest that individuals with higher fat intakes are more likely to develop disturbances of glucose metabolism, type 2 diabetes, and impaired glucose tolerance than individuals consuming lower amounts of fat, although obesity and physical inactivity may be confounding factors (215). Studies in experimental ani-
mals indicate that high-fat feeding induces both a decline in insulin sensitivity (216–218) and the development of insulin resistance in a variety of tissues (219–225).

1. Level of Dietary Fat

Cellular mechanisms responsible for the decline in insulin responsiveness in association with high-fat feeding have not been fully defined. Euglycemic, hyperinsulinemic clamp studies in human subjects and experimental animals indicate that high-fat feeding significantly impairs insulin action (226), reduces skeletal muscle glucose metabolism (226), and decreases the ability of insulin to suppress hepatic glucose production (226,227). Such studies also indicate that diet-associated development of peripheral insulin resistance may be modulated by age (228).

Reductions in insulin binding in tissues from rats fed high-fat as compared with high-carbohydrate diets have been observed by several groups of investigators (219,229–232). However, other investigators failed to observe significant alterations in insulin binding in response to high-fat feeding (220,233,234). Several postreceptor defects in insulin action (219,220,222,233–236) are observed in tissues from animals fed high-fat diets. Specifically, reductions in insulin receptor kinase activity (234), in the intracellular glucose transport system (222,233,235), and in the intracellular capacity to utilize glucose for lipogenesis (220) have all been reported in association with high-fat feeding.

2. Type of Dietary Fat

Although all diets high in fat lead to insulin resistance relative to high-carbohydrate diets, the effect of dietary fat on insulin action is greatly influenced by the type of fatty acid consumed [see reviews by Storlien et al. (237) and Lovejoy (238)]. High saturated fat intakes are consistently associated with insulin resistance, whereas monounsaturated and polyunsaturated fatty acids are less deleterious in this regard (215,237,238). van Amelsvoort et al. (239) observed that insulin response is greater in epididymal fat cells from rats fed diets high in polyunsaturated as compared with saturated fats. Likewise, diets with increasing ratios of polyunsaturated to saturated fatty acids induce alterations in the composition of adipocyte plasma membranes that are associated in a dose-dependent manner with increases in insulin binding, insulin receptor signaling, and glucose transporter activity (240–243). In addition, several studies indicate that substitution of more saturated with less saturated dietary fat sources can improve or ameliorate high-fat diet–induced impairment in insulin function (225,239,244,245). Substitution of safflower oil (a polyunsaturated fat) for beef tallow (a saturated fat) in a moderate-fat diet leads to an increase in the glucose uptake response to insulin (225) and to alterations in gene expression of several insulin signal transduction pathway intermediates (246). Specifically, the abundance of insulin receptor substrate 1 (IRS-1) and phosphatidylinositol (PI) 3-kinase mRNA and protein are lower in rats fed a beef tallow as compared with a safflower oil diet (246). In contrast, insulin receptor mRNA, relative expression of insulin receptor mRNA isoforms, and receptor protein are not influenced by dietary fat type (246). Recent evidence demonstrating differing effects of individual fatty acids on insulin-stimulated transport/phosphorylation, but similar effects on insulin-stimulated glycogen synthesis indicates that dietary fat type may interact with various metabolic pathways to different degrees and through different mechanisms (247).
It has been suggested that n-3 polyunsaturated fatty acids may afford a protective effect against diet-induced insulin resistance (215,237,243). Replacement of a small portion (6–11%) of the fatty acids in a high-safflower oil diet with long chain n-3 fatty acids from fish oil prevents the accumulation of intramuscular triglyceride and development of insulin resistance typically observed in association with consumption of a very high (59% calories) safflower oil diet (244,245). In insulin-resistant slightly diabetic rats, 6 weeks of fish oil feeding corrected the inhibitory effects of high-sucrose/high-fat feeding on insulin action (248). In that study, adipocytes from the fish oil–fed rats, as compared with those of corn oil–fed rats, had significantly higher rates of insulin-stimulated glucose transport, oxidation, and incorporation into total lipids that were positively correlated with the fatty acid unsaturation index of adipocyte membrane phospholipids. More recently, Podolin et al. (243) reported that the insulin resistance typically induced in rats by high-sucrose feeding could be prevented by adding as little as 6% fish oil to the diet. Recent data from human studies indicate that the degree of insulin resistance is negatively correlated to the amount of 22:6n-3 within skeletal muscle phospholipids (249). However, long-term and beneficial effects of n-3 fatty acids on insulin action are not consistently observed. Ezaki et al. (250) reported a stepwise increase in both insulin-stimulated glucose uptake and glucose transporter distribution following short-term substitution (1-week) of a portion of the safflower oil in a high-fat diet with fish oil. However, this effect is transient as further fish oil feeding (4 weeks) results in levels of insulin resistance and adipose cell enlargement comparable to those observed with the high-safflower diet. In addition, some human studies suggest that fish oil ingestion may exacerbate existing type 2 diabetes (251).

3. Reversibility of Diet-Induced Alterations in Insulin Action

Prolonged consumption of a high-fat diet impairs insulin action and leads to the development of obesity. Conversely, a reduction in dietary fat content may improve insulin sensitivity and reduce obesity development. Harris and Kor (217) observed an impaired insulin response to a glucose challenge in rats fed high-fat (40% energy) diets for 8 weeks. This effect is reversed within 3 days following a modest reduction in dietary fat content (30% energy). As alterations in body weight or fat content are not observed until 14 days following the switch to the lower fat diet, the improvement in insulin sensitivity is not believed to be secondary to a reduction in obesity (217). Though the above results are encouraging, an improvement in insulin response upon reduction in dietary fat content has not been consistently observed. In rats previously fed high-fat (60% energy) diets for 6 months, Yakubu et al. (218) failed to detect either an improvement in insulin response or a reduction in body weight following subsequent consumption of a low-fat (20% energy) diet for 3 or 6 months. Thus, alterations in insulin response induced by a high-fat diet and the reversibility of these effects appear to be influenced by both level and type of dietary fat as well as the duration of the high-fat feeding.

V. DIETARY FAT AND ADIPOSE TISSUE CELLULARITY

The expansion of adipose tissue during growth or the development of obesity is achieved through an increase in adipose tissue size (cellular hypertrophy), an increase in adipose tissue number (cellular hyperplasia), or through a combination of both
processes. Numerous studies provide evidence that variations in the level and type of fat included in the diet can lead to alterations in adipose cell size and/or number. As quantification of total adipose tissue cell number in human subjects is not readily obtainable, these studies have been primarily conducted in experimental animals. The major investigations concerning the influence of dietary fat level and type on adipose tissue cellularity are presented below.

A. Level of Dietary Fat

Many animal studies confirm that high-fat feeding leads to an expansion of adipose tissue mass through an increase in fat cell size and/or number and to the subsequent development of obesity. Studies of adipose tissue development in rodents indicate that increases in fat pad weight are typically associated with increases in both fat cell size and number until approximately 10–18 weeks of age (252–254). Body and fat pad weight and fat cell size and number then plateau and remain fairly constant in the adult animal. High-fat feeding influences both the dynamic stage of adipose tissue development early in life (252,255,256) and also the more static phase associated with adulthood (252,257,258). In mice fed high-fat diets from birth, increased fat pad weights are associated with a greater fat cell size through 18 weeks of age; this is followed by an increase in fat cell number through 52 weeks of age (252). At that time fat pad weight is sixfold greater in the high-fat compared with the control-fed mice, while fat cell size and number are increased 2.3- and 2.5-fold, respectively. An even greater effect of high-fat diets on adipose tissue cellularity is observed in young Osborne–Mendel rats (255), a strain susceptible to high-fat feeding (50). In these rats, a 4- to 16-fold increase in adipocyte number (dependent on the specific fat pad studied) is observed between 24 and 105 days of age when the animals are fed a high-fat diet (255).

Dramatic alterations in adipose tissue cellularity are also observed in adult rats subjected to high-fat feeding (252,257,258). In adult rats, high-fat diet–induced obesity is associated with increases in both fat cell size and fat cell number, with increases in cell size preceding changes in cell number. In 5-month-old rats, a significant increase in fat cell size is detected as early as 1 week after the introduction of a high-fat diet (257). This response is followed by increases in cell number in the perirenal fat pad after 2 weeks of diet treatment and in the epididymal fat pad after 8 weeks on the high-fat diet. Likewise, Faust et al. (258) report increases in cell size in several fat pad depots after 3 weeks of high-fat feeding in adult rats but an increase in cell number only after 9 weeks of dietary treatment. Fat cell hyperplasia in response to high-fat feeding is also observed in adult genetically obese rodents, with the magnitude of response being depot-dependent (259).

B. Type of Dietary Fat

It is generally accepted that a high level of fat in the diet may induce adipose cell hypertrophy and/or hyperplasia. However, the influence of dietary fat type on adipose tissue cellularity and the development of obesity is less definitive, particularly with respect to diet-induced alterations in fat cell hyperplasia. Several studies reported a greater effect of unsaturated as compared with saturated fat diets on increasing fat cell number (260–262). Specifically, an increase in the DNA content of the epididymal fat pad (indicative of an increased cell number) is observed in growing rats
fed diets containing 20% sunflower oil, whereas an increase in fat cell size is observed in those fed diets containing 20% lard. It is suggested (260) that the multiplication rate of adipose tissue cells may be increased when the degree of unsaturation of the dietary lipids is greater. However, this suggestion is contradicted by other investigations that report a greater degree of fat cell hyperplasia with saturated as compared with unsaturated fat diets (256,263,264). Lemmonier et al. (263) observed an increase in fat cell number in the perirenal depots of rats fed high levels of saturated fat and an increase in fat cell size in rats fed an unsaturated fat diet. In the epididymal fat pad, a similar degree of hypertrophy (and no cell hyperplasia) is observed in rats fed either of the high-fat diets compared with the low-fat diet controls (263). Bourgeois et al. (256) also reported adipose tissue site and sex-specific responses to various types of high-fat diets, with the greatest increases in fat cell number observed in those animals fed the saturated fats. In contrast, several studies suggested that the alterations in fat cell size and number associated with high-fat feeding are not influenced by type of dietary fat (239,265–267). Thus, Kirtland et al. (265) observed that long-term feeding of high-fat diets as compared with low-fat diets (20% and 3% w/w, respectively) in guinea pigs results in an increase in fat pad weight and fat cell size. However, quite similar results are observed with inclusion of either beef tallow or corn oil in the high-fat diet formulation. Likewise, nonobese mice respond to high-fat diets containing either corn oil or tallow with similar increases in body and fat pad weight and adipocyte volume over the low-fat diet controls (266).

In contrast to the studies reported above, more consistent effects on adipose cellularity are observed with dietary n-3 polyunsaturated fatty acids which selectively limit hypertrophy and/or hyperplasia in male rats in a depot-dependent manner (153,268). Whether differences in fat cell size and/or number are observed appears to be dependent on the duration of the study. Cell size is decreased during short-term n-3 fatty acid feeding studies (153,154,269,270), whereas fat cell number is also reduced during longer term studies in male (268) but not female (Higbee et al. unpublished observations) rats. Hill et al. (268) reported that retroperitoneal fat pads are most responsive to dietary manipulation followed by epididymal pads, whereas subcutaneous (inguinal) adipose tissue is slowest to respond. They observed a reduction in both fat cell size and number in retroperitoneal fat of male rats with fish oil feeding (as compared with lard or corn oil) after 3 months. Similar reductions in adiposity were observed in epididymal and inguinal depots, although only cell size was affected. In contrast, Higbee et al. (unpublished observations) observed only minor alterations in cell size distribution and no significant differences in fat cell number of inguinal, retroperitoneal, or parametrial fat depots between female rats fed high-fat diets containing either fish oil or corn oil for up to 32 weeks. These discrepancies suggest possible gender differences in response to dietary fats that warrant further investigation.

C. Interrelationship Between Changes in Fat Cell Size and Number

As previously mentioned, studies of the influence of the level or type of dietary fat on adipose tissue cellularity frequently observe site- and sex-specific differences in response (252,256,264). When adipose tissue development in animals with genetic or diet-induced obesity is monitored over time, it is generally observed that increases in fat cell number and size are initially observed in the retroperitoneal (252,264,266,269,270) but
monier (252) suggested that in certain conditions there might be a “maximum cell size” which is similar regardless of animal sex or adipose depot site. Thus, the greater degree of hyperplasia observed in the perirenal compared with other fat depots in response to high-fat feeding may be due to the fact that cells of this depot are already near-maximal in size [for review, see (259)]. Likewise, Faust et al. (258) advanced the notion that during the development of obesity in adult rats fed a high-fat diet, the achievement of some specific mean fat cell size triggers the events that lead to an increase in adipocyte number. They further suggested that when this “critical” fat cell size is reached, a stimulus for new cell production or differentiation may be produced.

Several studies indicated that the feeding of a high-fat diet to adult rats leads to an increase in the incorporation of [³H]thymidine into the DNA of specific adipose tissue fractions (271,272), providing direct evidence that the diet-induced increase in fat cell number is indeed a consequence of cellular proliferation. However, a direct link between an increase in cellular proliferation and the release of factors from adipose tissue triggering this effect has been difficult to demonstrate. For example, Bjorntorp et al. (273) reported that addition of plasma from either chow-fed or high-fat-fed rats to adipocyte precursors in culture has similar effects on inducing preadipocyte proliferation and differentiation. Likewise, Shillabeer et al. (274) and Lau et al. (275) observed that media conditioned by exposure to mature fat or cells obtained from adipose tissue influence the proliferation and differentiation of cultured preadipocytes. However, a similar effect on preadipocyte differentiation is observed with fat from rats fed differing levels and types of dietary fat (264). In contrast, compelling evidence for such a relationship between diet-induced alterations in adipose tissue cellularity and the release or modification of local growth factors from adipose tissue of high-fat diet–treated animals was obtained in recent studies from our laboratory (276,277). These studies indicated that changes in adipose cellularity during the development of obesity in rats fed high-fat diets may be associated with the appearance of a locally produced factor(s) capable of stimulating adipose cell proliferation. Specifically, we observed that rats fed high-fat diets have increased numbers of both very small and large cells within their adipose tissue depots and that media conditioned by exposure to the fat from these animals significantly increase the proliferation of preadipocytes in primary culture (276). In addition, we determined that total tissue content of insulin-like growth factor type I (IGF-I) is greater in the fat pads of high-fat diet rats as compared with their low-fat diet controls (277). Furthermore, adipogenic activity in the conditioned media from the high-fat diet rats could be attenuated through neutralization with an IGF-I antibody (276). Taken together, these studies support the concept that the enlargement of adipocytes during the development of diet-induced obesity may lead to the secretion of local growth factors involved in the regulation of adipose tissue expansion and that IGF-I may be at least partially responsible for this effect (Fig. 2).

D. Cellularity Changes and the Reversibility of Obesity

The potential reversibility of obesity induced by high-fat feeding may be dependent on the duration and/or severity of the dietary treatment (258,278,279). For example, diet-induced obesity is reversed when the animals are returned to a low-fat diet after 16 (278) but not after 30 weeks (279) of high-fat feeding. It has been proposed that
the specific diet-induced alterations in adipose tissue cellularity may be a major factor influencing the reversal of the obese state (258,278). Thus, obesity associated with changes in fat cell size is readily reversed only when the animals are returned to a low-fat regime (258,278). However, this is not the case for obesity associated with changes in fat cell number. Diet-induced changes in fat cell number appear to be permanent, as substitution of a high-fat diet with a low-fat diet leads to reductions in body weight and fat cell size but not in fat cell number (258,278,279). Thus, a permanent increase in the capacity for energy storage and in the potential for expansion of adipose tissue mass would be a consequence of long-term exposure to the high-fat feeding. As mentioned previously (Sec. II), high levels of dietary fat have been associated with the development of obesity in humans. If the observations regarding cellularity alterations and the potential reversal of obesity are indeed applicable to humans, this would suggest that high-fat diet–induced obesity, particularly that of extended duration, may be resistant to intervention and may also have long-term metabolic consequences.

REFERENCES

7. B. M. Popkin, S. Paeratakul, F. Zhai, and K. Ge. A review of dietary and environmental 
9. Z. S. Warwick and S. S. Schiffman. Role of dietary fat in calorie intake and weight 
11. A. Kendall, D. A. Levitsky, B. J. Strupp, and L. Lissner. Weight loss on a low-fat diet: 
    Cole, and L. Brace. Changes in body weight, body composition and energy intake in 
15. J. E. Blundell, C. L. Lawton, J. R. Cotton, and J. I. Macdiarmid. Control of human 
    B. McCormick, D. M. Bier, and A. G. Goodridge, eds.). Annual Reviews, Palo Alto, 
16. C. P. Sepple and N. W. Read. The effect of pre-feeding lipid on energy intake from a 
17. L. L. Birch, L. S. McPhee, J. L. Bryant, and S. L. Johnson. Children’s lunch intake: 
    Effects of mid-morning snacks varying in energy density and fat content. *Appetite* 20: 
    83 (1993).
18. C. de Graaf, T. Hulshol, J. A. Westrate and P. Jas. Short term effects of different 
19. J. R. Cotton, V. J. Burley, J. A. Westrate, and J. E. Blundell. Dietary fat and appetite: 
    Similarities and differences in the satiating effect of meals supplemented with either 
    control in obese subjects: Weak effects on satiation and satiety. *Int. J. Obes. Relat. 
    Metab. Disord.* 17:409 (1993).
    T. Houtsmuller. Effects of varying the carbohydrate:fat ratio in a hot lunch on post- 
22. S. M. Green, V. J. Burley, and J. E. Blundell. Effect of fat- and sucrose-containing 
    foods on the size of eating episodes and energy intake in lean males: potential for 
    of energy intake: Evaluating the effects of fat on meal size and post-meal satiety. *Am. 
    Stoner. Satiety after preloads with different amounts of fat and carbohydrate: Implica- 


Lipid-Based Synthetic Fat Substitutes

CASIMIR C. AKOH

The University of Georgia, Athens, Georgia

I. INTRODUCTION: WHY SYNTHETIC FAT SUBSTITUTES?

Fat is an important macromolecule component of plant and animal tissues. The various fats contribute to the physical and functional properties (solubility, viscosity, rheology, melting behavior, emulsification, body, creaminess, heat conduction, carrier of lipophilic vitamins and flavorants) of most food products, affecting as well sensory (appearance, taste, mouthfeel, lubricity, flavor) and nutritional (satiety, calories, essential fatty acid source, health benefits) aspects of food. These properties are difficult to duplicate in food formulations without adding fats. The amount and type of fat present in a food determine the characteristics of that food and consumer acceptance.

Fat is still the number one nutritional concern for most people in developed countries. The present estimate of average fat calories consumed by most Americans is 35–37%. Recommendation by the U.S. Senate Select Committee on Nutrition and Human Needs and the Surgeon General is that fat consumption be reduced to 30% of total calories of the diet (1). Excessive intake of fat in the diet has been linked to certain diseases, such as heart disease, cancer, obesity, and possibly gallbladder disease (2,3). Increased saturated fat intake is associated with high blood cholesterol and increased risk of coronary heart disease. It has been difficult for individuals to change their dietary habits to reduce or minimize fat intake while enjoying their favorite foods. This problem and the interest shown by consumers in alternative fats and foods low in calories or without calories led to the search by the food industry and scientific community for the “ideal” fat substitute (4,5).

A real fat substitute must be able to provide all the attributes of fats and replace the calories from fat on a 1:1 weight basis (2). Fats are the most concentrated source
of energy: a given amount of fat contains more than twice the calories (9 kcal/g) of other macronutrients such as proteins and carbohydrates (4 kcal/g). An “ideal” fat substitute must look and function like fat and be able to substantially reduce caloric contributions to food. In recent years, health-conscious consumers have shown interest in reducing calories from fat by modifying their diets, exercising, and eating healthier foods. Recent survey results show that consumers are ready for fat substitutes and replacers in their foods (6).

II. CLASSIFICATION OF FAT REPLACERS

Fat replacers are divided into two main groups: fat mimetics and fat substitutes. They are classified as carbohydrate-based, protein-based, and lipid-based fat replacers (Table 1), or combinations thereof. The protein-based and carbohydrate-based replacers are widely regarded as fat mimetics. The fat mimetics are either proteins or carbohydrates that have been physically or chemically processed to mimic the properties and functions of fats in food systems; they are not fats. They tend to adsorb a large amount of water, are not stable at frying temperature, and may produce food that is not microbiologically shelf stable; they contribute some calories (1–4 kcal/g) to the diet. The fat mimetics do not possess all the organoleptic, physical, chemical, and functional properties of fats and cannot replace calories from fat on a 1:1 weight basis. They cannot carry lipid-soluble flavor compounds because they cannot lower the vapor pressure of lipophilic flavor molecules, and most foods prepared with fat mimetics are often perceived by consumers as lacking in taste. Although fat mimetics need a delivery system such as an emulsifier to carry lipid-soluble flavors, they can carry water-soluble flavors. Some fat mimetics have mouthfeel and physical properties approximating those of triacylglycerols but are not suitable for frying operations because they can be denatured (protein-based substances) or caramelized (those based on carbohydrates). They can, however, be used for baking and retort cooking operations.

Fat substitutes are believed to be compounds that physically and chemically resemble triacylglycerols. They are stable to high temperature cooking and frying operations.

### Table 1 Typical Examples of Fat Replacers

<table>
<thead>
<tr>
<th>Compound</th>
<th>Class</th>
<th>Caloric content (kcal/g)</th>
<th>Absorbability</th>
<th>Uses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Olestra/Olean</td>
<td>Lipid-based</td>
<td>0</td>
<td>Nonabsorbable</td>
<td>FDA approved for savory snacks. Stable to frying, baking, and cooking temperatures. Supplementation with fat-soluble vitamins required. Can be used for dairy products, spreads, and dressing but will require separate approval.</td>
</tr>
</tbody>
</table>
operations, and in theory, can replace fats and oils on a 1:1 weight basis in foods. In the literature, the term “fat substitutes” has been used interchangeably with “fat replacers,” but not with “fat mimetics,” and this can be confusing. Several lipid-based synthetic low-calorie or zero-calorie fat substitutes belong to the fat substitute group. A good example is sucrose fatty acid polyester (SPE), which was originally developed as olestra and marketed as Olean by Procter & Gamble to replace edible fats and oils in the diet (7). A number of other fat substitutes have been developed or are under development (8–18). Among these, the carbohydrate and alkyl glycoside fatty acid polyesters and the structured lipids (SLs) have functional and physical properties resembling those of triacylglycerols while contributing few to no calories to the diet (10–20).

Some of the lipid-based fat substitutes can be added to food products to replace the functional properties of fats, including frying (not possible with protein-based fat replacers such as Simplesse®, and carbohydrate-based fat replacers such as maltodextrin), while reducing caloric contributions from fats and oils. Table 2 lists the applications and functions of some fat replacers. Because the “ideal” fat substitute does not exist, a systems approach to reduced-fat or low-fat food formulations has been proposed. Each type of food product will require a different approach to address the difficulties of formulating a counterpart that has reduced, no, or low fat. Simply put, a systems approach uses a combination of different ingredients that may or may not belong to either of the classes of fat replacers and requires a basic knowledge of ingredient technology to formulate desired products. The system may contain emulsifiers, fat substitutes or mimetics, fibers, water control ingredients, flavor, and bulking agents. Water or moisture control poses one of the greatest challenges in formulating reduced-fat snack and baked goods. In these systems water is used to replace fat, to increase bulk, or for functionality. A detailed review of fat mimetics is outside the scope of this chapter, which concentrates on lipid-based fat substitutes.

III. TYPES OF LIPID-BASED SYNTHETIC FAT SUBSTITUTES

Lipid-based fat substitutes include carbohydrate fatty acid polyesters such as sucrose polyester, sorbitol polyester, raffinose polyester, stachyose polyester, and alkyl glycoside fatty acid polyesters. Others include Caprenin, Salatrin (short and long acyl triglyceride molecules, marketed as Benefat), structured lipids, medium chain triacylglycerols (MCTs), mono- and diacylglycerols, esterified propoxylated glycerol (EPG), dialkyl dihexadecylmalonate (DDM), and trialkoxytricarballylate (TATCA), to name a few. The composition and sources or developers of the lipid-based fat substitutes are shown in Table 3.

Of all the lipid-based fat substitutes, only sorbitol, trehalose, raffinose, and stachyose polyesters have a chance to compete with Olestra as nondigestible zero-calorie fat substitutes. Others are either partially hydrolyzed or fully hydrolyzed and absorbed, thus contributing some calories to the diet.

A. Strategies for Designing Lipid-Based Fat Substitutes

Several strategies were suggested for designing low-calorie or zero-calorie lipid-based synthetic fat substitutes (2,16). The basic premise is to reengineer, redesign, chemically alter, or synthesize conventional fats and oils such that they retain the
Table 2  Applications and Functions of Some Fat Replacers

<table>
<thead>
<tr>
<th>Specific application</th>
<th>Fat replacer</th>
<th>General functions&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baked goods</td>
<td>Lipid-based</td>
<td>Emulsification; cohesiveness; tenderizer, flavor carrier, shortening replacer, antistaling agent; prevention of retrogradation of starch; dough conditioner</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Moisture retention; retard staling</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Texturizer</td>
</tr>
<tr>
<td>Frying and cooking</td>
<td>Lipid-based</td>
<td>Texturizer; flavor, crispiness; heat conduction</td>
</tr>
<tr>
<td>Salad dressing</td>
<td>Lipid-based</td>
<td>Emulsification; mouthfeel; hold flavorants</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Increase viscosity; mouthfeel; texturizer</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Texturizer; mouthfeel</td>
</tr>
<tr>
<td>Frozen desserts</td>
<td>Lipid-based</td>
<td>Emulsification; texture</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Increase viscosity; texturizer, thickener</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Texturizer, stabilizer</td>
</tr>
<tr>
<td>Margarines, shortenings, spreads, and butter</td>
<td>Lipid-based</td>
<td>Spreadability; emulsification; flavor; plasticity</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Mouthfeel</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Texturizer</td>
</tr>
<tr>
<td>Confectionery</td>
<td>Lipid-based</td>
<td>Emulsification; texturizer</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Mouthfeel; texturizer</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Mouthfeel; texturizer</td>
</tr>
<tr>
<td>Processed meat products</td>
<td>Lipid-based</td>
<td>Emulsification; texturizer; mouthfeel</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Increase water-holding capacity; texturizer; mouthfeel</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Texturizer; mouthfeel; water holding</td>
</tr>
<tr>
<td>Dairy products</td>
<td>Lipid-based</td>
<td>Flavor; body; mouthfeel; texture; stabilizer; increase overrun</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Increase viscosity; thickener; gelling agent; stabilizer</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Stabilizer; emulsification</td>
</tr>
<tr>
<td>Soups, sauces, and gravies</td>
<td>Lipid-based</td>
<td>Mouthfeel; lubricity</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Thickener; mouthfeel; texturizer</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Texturizer</td>
</tr>
<tr>
<td>Snack products</td>
<td>Lipid-based</td>
<td>Emulsification; flavor</td>
</tr>
<tr>
<td></td>
<td>Carbohydrate-based</td>
<td>Texturizer; formulation aid</td>
</tr>
<tr>
<td></td>
<td>Protein-based</td>
<td>Texturizer</td>
</tr>
</tbody>
</table>

<sup>a</sup>Functions are in addition to serving as a fat replacer.

conventional functional and physical properties of fats and oils in foods but contribute few or no calories because of reduced susceptibility to hydrolysis and/or absorption in the lumen. Possible strategies, rationale, and examples are as follows:

1. Replace the glycerol moiety of the triacylglycerol with alternative alcohols (e.g., carbohydrates, polyols, neopentyl alcohol). This ensures steric protection of the ester bonds. Branching interferes with hydrolysis by pancreatic lipase. Examples are sucrose fatty acid esters, sucrose polyesters, other
Table 3  Types of Lipid-Based Fat Substitutes

<table>
<thead>
<tr>
<th>Name</th>
<th>Composition</th>
<th>Source/developer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Olestra or Olean</td>
<td>Sucrose polyester of fatty acids (6–8 fatty acids)</td>
<td>Procter &amp; Gamble (FDA-approved, 1996), Akoh and Swanson, Unilever</td>
</tr>
<tr>
<td>Caprenin</td>
<td>Caprocaprylohebenin-structured triacylglycerol C8:0, C10:0, C22:0</td>
<td>Procter &amp; Gamble. GRAS requested^</td>
</tr>
<tr>
<td>Salatrim/Benefat</td>
<td>(C18:0, C2:0, to C4:0)-structured triacylglycerol</td>
<td>Nabisco Foods Group/Cultor Food Science</td>
</tr>
<tr>
<td>EPG</td>
<td>Esterified propoxylated glycerols</td>
<td>ARCO Chemical Co. CPC International</td>
</tr>
<tr>
<td>DDM</td>
<td>Dialkyl dihexadecylmalonate</td>
<td>Frito-Lay, Inc.</td>
</tr>
<tr>
<td>TATCA</td>
<td>Trialkoxytricarboxylate</td>
<td>CPC International</td>
</tr>
<tr>
<td>TAC</td>
<td>Trialkoxyxycitrate</td>
<td>CPC International</td>
</tr>
<tr>
<td>Alkyl glycoside polyesters</td>
<td>Alkyl glycosides + fatty acids</td>
<td>Akoh and Swanson, Curtice Burns, Inc.</td>
</tr>
<tr>
<td>Trehalose, raffinose, stachyose polyesters</td>
<td>Carbohydrate + fatty acids (all similar to olestra)</td>
<td>Akoh and Swanson, Curtice Burns, Inc.</td>
</tr>
<tr>
<td>Sorbestrin</td>
<td>Sorbitol or cyclic sorbitol + fatty acids</td>
<td>Cultor Food Science</td>
</tr>
<tr>
<td>PGE</td>
<td>Polglycerol esters–emulsifiers</td>
<td>Lonza, Inc.</td>
</tr>
<tr>
<td>Sucrose esters</td>
<td>Sucrose with 1–4 fatty acids as emulsifiers</td>
<td>Mitsubishi Chemical America, Inc., Crodesta</td>
</tr>
<tr>
<td>TGE</td>
<td>Trialkoxyglycerol ether</td>
<td>CPC International</td>
</tr>
<tr>
<td>MCT</td>
<td>Medium chain triglycerides</td>
<td>ABITEC Corp., Stepan Co.</td>
</tr>
<tr>
<td>Phenylmethylpolysiloxane</td>
<td>Organic derivatives of silica</td>
<td>Dow Corning Corp.</td>
</tr>
</tbody>
</table>

^Developer has petitioned the FDA to obtain “Generally Recognized as Safe” status for the product.

carbohydrate fatty acid polyesters, alkyl glycoside fatty acid polyesters, and polglycerol esters.

2. Replace the long chain fatty acids with alternative acids (to confer sterific protection to the ester bonds). Examples are branched carboxylic esters of glycerol and structured lipids such as Caprenin and Salatrim/Benefat. Caprenin and Salatrim contain poorly or less absorbed long chain saturated fatty acids and easily absorbed short and/or medium chain fatty acids esterified to the glycerol. The short and medium chain fatty acids have lower heats of combustion than long chain fatty acids.

3. Reverse the ester linkage in triacylglycerols by replacing the glycerol moiety with a polycarboxylic acid, amino acid, or other polyfunctional acid and esterify with a long chain alcohol. Examples include trialkoxytricarboxylate (TATCA) and trialkoxyxycitrate (TAC).

4. Reduce the ester linkage of the glycerol moiety to an ether linkage. This product is not a good substrate for lipases, which do not hydrolyze ether.
bonds as fast as ester bonds. Examples include diether monoesters of glycerol, triglyceryl ethers, and trialkoxyglyceryl ether (TGE).

5. Apply chemistry unrelated to triacylglycerol structure. A good example is the use of polymeric materials having physical and functional properties similar to those of conventional fats and oils such as phenylmethylsiloxane (PS) or silicone oil and paraffins.

6. Evaluate naturally occurring substances as potential low-calorie fat substitutes. Jojoba oil is an excellent example.

7. Use enzymes to synthesize reduced-calorie fat substitutes. Examples include sugar mono- and diesters, glycerophospholipids, mono- and diacylglycerols, and structured lipids.

8. Introduce oxypropylene group between glycerol and fatty acids to form propoxylated molecules. An example is esterified propoxylated glycerol (EPG).

B. Olestra or Sucrose Polyester: Brief History of Development

Sucrose fatty acid polyester (SPE) development dates back to the year 1880, when a derivative of sucrose was prepared by acetylation to produce sucrose octaacetate (i.e., sucrose containing eight acetate groups). Following this, other carbohydrate acetates were successfully prepared. In 1921 Hess and Messner (21) synthesized sucrose octapalmitate (sucrose esterified with eight molecules of palmitic acid, a long chain fatty acid) and sucrose octastearate. In 1952 the concept of sucrose polyester production was initiated when the president of the Sugar Research Foundation, Henry B. Hass, asked Foster D. Snell to look into the possibility of “hanging a fat tail on sucrose” for use in detergents. The idea was that since sucrose is highly hydrophilic, a lipophilic tail on sucrose would result in a molecule that is amphiphilic (both water and oil loving), hence able to serve as an excellent surfactant. It was anticipated that production would be easy and the product biodegradable under aerobic and anaerobic conditions. It turned out that the chemical synthesis was not that easy without the use of solvents like dimethylformamide (DMF), dimethylsulfoxide (DMS), and dimethylpyrrolidone (DMP) to solubilize sucrose and free fatty acids. This process, called the Hass-Snell process, was applicable only to the synthesis of sucrose mono- and diesters, otherwise called sucrose fatty acid esters (SFEs). These are digestible and good nonionic surfactants, as we shall see later in this chapter. By today’s standard, the solvents used are not food grade; therefore, products made in them are unacceptable for human consumption.

The other concept was to find means of reducing fat-derived calories without resorting to dilution with, say, water, air, carbohydrates, and proteins. The aim was to somehow come up with a fatlike molecule that would significantly reduce fat calories by preventing their hydrolysis and absorption. This led to the discovery of a nondigestible and nonabsorbable fatlike molecule called sucrose fatty acid polyester, now known as olestra (generic name) or Olean (brand name), by Mattson and Volpenhein (7) while working on the absorption of fats by infants.

Sucrose is a nonreducing disaccharide and the common table sugar. “Olestra” or “sucrose polyester” refers to sucrose esterified with six to eight fatty acids. SPEs become undigestible when the number of fatty acids esterified is greater than 4. The structure of sucrose polyester is given in Figure 1. Procter & Gamble, which was
granted the original patent for sucrose polyester in 1971, spent over $250 million over the last 25 years to develop this fat substitute. The original application for use of sucrose polyester as a food additive, filed with the Food and Drug Administration (FDA) in April 1987, was withdrawn and modified. But on January 24, 1996, the FDA approved olestra for limited use in savory snacks (namely, chips, curls, and crackers). Before the approval, sucrose polyester was evaluated in over 10 animal studies and in 25–30 clinical trials. The approval of olestra was not without controversy. The Center for Science in the Public Interest (CSPI), a Washington consumer advocate group, believes that olestra deprives the body of some of the essential vitamins and carotenoids that may protect against cancer. In 1996, in Iowa, Wisconsin, and Colorado, Frito-Lay, a unit of PepsiCo, test-marketed chips made with the olestra.

1. Synthetic Approaches

Sucrose fatty acid polyester can be synthesized in the presence or absence of organic solvent. Direct esterification of sucrose with fatty acid is very difficult. The solvent-free process is widely used for the current production of sucrose polyester (18,22). The synthesis may involve reactions of the following types:

1. **Transesterification.** Fatty acid methyl esters (FAMEs) and sucrose are reacted in the presence of potassium soaps to form a homogeneous melt followed by the addition of excess FAME and NaH at 130–150°C. In some cases, potassium carbonate is added to aid the reaction. The function of soap is to help solubilize sucrose and FAME. Methanol, a by-product of the transesterification reaction, is distilled off (Fig. 2). The active catalyst is the sucrate ion generated with alkali metal hydrides. This is a two-stage transesterification process, and up to 8–9 hours may be needed to achieve 90% yield of SPE.

2. **Interesterification (ester interchange).** This involves reacting a short chain alkyl ester such as sucrose octaacetate (SOAc) with FAME in the presence of sodium methoxide (NaOH) or Na or K metal as catalyst (14). The reaction requires extremely anhydrous conditions to prevent hydrolysis of formed product, catalyst inactivation, or explosion (when Na is in contact with water). The temperature of the reaction with Na catalyst is lower.
Figure 2 Synthetic scheme for olestra by transesterification and processing.

(105–130°C). Reaction times of 2–6 hours and pressure of 0–5 mm Hg are required to achieve greater than 95% yield of SPE (14). This is a simple ester–ester interchange reaction (Fig. 3). The methyl acetate formed is trapped with a liquid nitrogen (−196°C).

Synthesis based on Na metal (potentially explosive and flammable compound) as catalyst and sucrose octaacetate and FAME as substrates may not be suitable for industrial adaptation. Therefore, use of milder catalysts such as sodium methoxide, potassium soap, and potassium carbonate are encouraged. Shieh et al. (23) recently reported the optimized synthesis of SPE using potassium hydroxide (KOH) in methanol plus FAME to form soap (potassium soap) followed by the addition of potassium carbonate. The reaction time at 144°C was 11.5 hours.

The triacylglycerol for the synthesis can come from vegetable oils and fats, alone or in combination. The type of product desired dictates the type of fatty acids needed for synthesis to achieve desired functionality. In most cases, the fatty acid profiles of finished products will resemble those of the triacylglycerol source (22). It should be noted that the reactions above are random processes and therefore the specific position and type of fatty acid on the sucrose molecule will vary from product to product.

2. Analyses of Olestra (SPE)

No matter what synthetic approach is used, at the end of the reaction, the vessel is cooled down to approximately 50°C or less and the product neutralized with acid,
washed, bleached, distilled (to remove unreacted substrates and sucrose ester with low degree of esterification), deodorized, and analyzed for extent of esterification (22). Antioxidant(s) may or may not be added to prevent oxidation during storage (Figs. 2 and 3). Olestra can be analyzed by any of the following techniques or combinations thereof:

1. Thin-layer chromatography (TLC), Iatroscan, and TLC-recording densitometry for degree of esterification (18).
2. Gas–liquid chromatography to determine the fatty acid composition of the product after transmethylation of olestra (10,18,24) or to determine olestra content after derivatization (silyl or acetate derivatives).
3. Column chromatography to separate products of small-scale synthesis (10,14).
4. High-performance liquid chromatography (HPLC) to analyze the content of olestra in, say, salad dressing, cooking oil, margarine, and spreads whose ingredients include triacylglycerols and fat-soluble vitamins, or analysis by size exclusion chromatography (SEC) for oxidized, polymerized, or heated olestra (25–27).
5. Fourier transform infrared spectroscopy (FTIR) for functional group and hydroxyl value determinations (14,18).
6. Nuclear magnetic resonance (NMR) spectroscopy for structural elucidation (14,18).
7. Supercritical fluid chromatography (SFC) of trimethylsilyl ether derivatives of olestra for separation and determination of degree of esterification (28).
8. Spectrophotometry for analysis of olestra content in foods (29).
10. Desorption mass spectrometry (MS) for characterization of olestra (31).
11. Hyphenated chromatography techniques, such as HPLC-FTIR, GC-MS, and HPLC-GC-FTIR, may also be useful.

Figure 4 shows the result of HPLC separation of olestra or sucrose polyester and triacylglycerol blend on a gel permeation chromatography column (32).

C. Olestra-Type Fat Substitutes

1. Sorbitol Polyester

Sorbitol or glucitol is a sugar alcohol (polyol) made by hydrogenation or electrolytic reduction of glucose. Other polyols like xylitol, mannitol, and lactitol can equally be used for the synthesis of fat substitutes. Up to six hydroxyl groups can be esterified with fatty acids to produce a nondigestible sorbitol polyester. The structure is shown in Figure 5. Sorbitol polyester can be synthesized by interesterification of sorbitol hexaacetate with FAME in the absence of organic solvent (13) or by transesterification reaction between sorbitol and FAME as described above.

Figure 4  HPLC chromatographic analysis of a mixture of olestra and triacylglycerol blend on four gel permeation chromatography columns arranged in series: a, olestra; b, a triacylglycerol; injection volume, 200 μL; flow rate, 1.0 mL/min; a light-scattering mass detector was used. (From Ref. 32.)
Recently, Chung et al. (33) optimized the synthesis of sorbitol fatty acid poly-
esters. Procter & Gamble also worked on sorbitol polyesters but did not follow up on their development. The properties and applications in food have not been exten-
sively studied as with olestra. However, sorbitol polyesters are proposed as zero-
calorie fat substitutes. They are less viscous than sucrose polyester and slightly more viscous than vegetable oils (13). Based on the known structure and limited metabolic studies, sorbitol polyesters may be used in place of olestra in food products. They are stable to high heat and taste and function like fats.

2. Sorbestrin

Sorbestrin is sorbitol or cyclic sorbitol containing three to five fatty acids esterified to the OH group. It was developed by Pfizer Inc. Sorbestrin contains 1.5 kcal/g and has a bland oil-like taste with cloud point between 15°C and 13°C. It is mainly a clear liquid and can serve as a reduced-calorie fat substitute. It is thermally stable and can withstand frying temperatures. It is intended for use in frying, baking, and salad dressings. Sorbestrin is not yet commercially available and will require food additive petition and FDA approval prior to use. Akoh and Swanson (13) synthesized sorbitol hexaoleate as a low-calorie fat substitute. Pfizer noted that for humans, sorbestrin consumption in the future is estimated at 12 g/day. The structure of sorbestrin is shown in Figure 6.

3. Trehalose Polyester

Trehalose is a nonreducing disaccharide, the major sugar of insect hemolymph, fungi, and yeasts. It is made of two α-D-glucopyranose components and closely resembles sucrose in physical properties. The nonreducing sugars are better substrates for the synthesis of fat substitutes than the reducing sugars. This is because the anomic carbon atoms (C-1) of nonreducing sugars are protected and not very susceptible to thermal degradation. The reducing sugars degrade and caramelize at the high temperatures required for transesterification. The use of acetyl derivatives of nonreducing sugars allows reactions to be carried out at reduced temperatures (<105–120°C). Unfortunately, this process is not applicable to acetylated reducing sugars such as glucose pentaacetate.
Trehalose octaacetate has been used to interesterify FAME catalyzed by Na metal to produce highly substituted trehalose polyester with properties similar to those of sucrose polyester (13). Therefore, trehalose polyester can be used as a zero-calorie fat substitute in place of sucrose polyester. The only problem is cost effectiveness, since sucrose is a cheaper substrate than trehalose. The structure of trehalose polyester is depicted in Figure 7.

4. Raffinose Polyester

Raffinose, like sucrose, is a heterogeneous nonreducing sugar. Raffinose is made of galactose–glucose–fructose units. It has 11 hydroxyl groups that can be esterified with fatty acids. It is well established that as the degree of substitution of a carbohydrate with fatty acids increases, the susceptibility to hydrolysis and absorption decreases. Based on this fact, raffinose undecaacetate was used to interesterify FAME to produce raffinose polyester containing 10–11 fatty acids in 99% yield. The reaction was catalyzed by Na metal at 110°C for 2–3 hours (10). The product had consistency in the range of salad oils and sucrose polyesters, with some raffinose polyesters being slightly more viscous than sucrose polyesters and salad oils depending on the fatty acid composition and degree of substitution. The structure of raffinose polyester is shown in Figure 8.
Metabolic studies in mice indicate that raffinose polyester is not hydrolyzed by intestinal lipases and is not absorbed (11). About 93–98% of the raffinose polyester fed to the mice was recovered unchanged from the feces. Thus, raffinose polyester can serve as an excellent zero-calorie fat substitute. Now that olestra has received FDA approval, more interest is expected on the possible use of raffinose polyester and others as alternative fat substitutes.

5. Stachyose Polyester

Stachyose, a nonreducing heterogeneous tetrasaccharide, is similar to raffinose and sucrose except that stachyose contains galactose–galactose–glucose–fructose. There are 14 available OH groups for esterification with long chain fatty acids. Stachyose can be completely acetylated and used as the substrate for interesterification with FAME, catalyzed by Na metal as described for sucrose and raffinose acetates. The degree of substitution as determined by $^{13}$C NMR spectroscopy was about 12 fatty acids per molecule of stachyose (12). The higher the degree of acetylation of the substrate, the lower the melting point, and the lower the temperature requirement for the interesterification. The only requirement is that the temperature be high enough to melt the Na catalyst (>98°C).

The structure of stachyose polyester is given in Figure 9. It is conceivable that this molecule will be highly resistant to hydrolysis and absorption and able to serve as an excellent zero-calorie fat substitute. The drawback is the cost, since again sucrose is cheaper.

D. Alkyl Glycoside Polyesters

Alkyl glycosides are prepared by reacting the sugar (e.g., glucose) with a desired alcohol (methanol for methyl glucoside) in the presence of acid catalysts. This glycosylation is important to convert the reducing C-1 anomerics centers to nonreducing, less reactive anomic centers. The alkyl glycosides become substrates for transesterification with FAME in the presence of potassium soap or sodium methoxide as catalysts. Alternatively, the remaining hydroxyl groups can be acetylated and used for the interesterification reaction catalyzed by Na metal as described for sucrose polyester, to produce alkyl glycoside fatty acid polyesters (12).
The different types of alkyl glycoside polyester synthesized by the latter process are shown in Figures 10–12. Up to 99% product yields have been reported. Physical properties are closer to those of vegetable oils than to sucrose polyester, partly because a maximum of only four fatty acids can be esterified. The alkyl glycosides are partially hydrolyzed and absorbed and can serve as reduced calorie but not as zero calorie fat substitutes.

E. Sucrose Esters (SFEs)

Sucrose fatty acid esters with a degree of substitution (DS) of 1–3 are highly hydrophilic, digestible, and absorbable; they are usable as solubilization, wetting, dispersion, emulsifying (especially oil-in-water, O/W, and some water-in-oil, W/O, emulsions), and stabilization agents, and as antimicrobial and protective coatings for fruits (8,34,35). They have a wide range of lipophilic–hydrophilic balance (HLB), namely, 1–16. They are tasteless, odorless, nontoxic, and biodegradable, and they can be used in food, cosmetic, and pharmaceutical applications. The structure of a sucrose monoester is shown in Figure 13.
Figure 11  Structure of methyl galactoside polyester.

Figure 12  Structure of octyl-β-glucoside polyester.

Figure 13  Structure of sucrose monoester, a sucrose ester.
1. Synthetic Approaches

Sucrose has eight hydroxyl groups that can be replaced or esterified with fatty acids. The problem for chemists was how to bring two immiscible molecules (sucrose and fatty acids or fatty acid methyl esters) to react in the presence of a suitable catalyst. A logical approach would be to dissolve them in a mutual solvent. To obtain a homogeneous solution of the reactants as required for synthesis of mono- and diesters of sucrose, a transesterification reaction was set up: that is, sucrose and free fatty acids were solubilized in mutual organic solvents such as dimethylformamide, dimethyl sulfoxide, and dimethylpyrrolidone. Sucrose fatty acid esters produced by this Hass–Snell process were not approved for use in food because of the potentially toxic solvents used (30,34). Later, Feuge et al. (36) described a solvent-free interesterification between molten sucrose and fatty acid methyl esters at 170–187°C in the presence of lithium, potassium, and sodium soaps as solubilizers and as catalysts. In both processes, the unreacted FAME and free fatty acids are removed by distillation. The higher the monoester content of the purified product, the better the SFEs are as emulsifiers. Enzymatic synthesis of SFE has not been very successful. Both buffer and organic solvent have been used to carry out enzymatic esterification of sucrose with mixed results (37).

2. Emulsification Properties

Sucrose fatty acid esters were approved for use in Japan in 1959 and in the United States in 1983 (34). Approved uses include:

1. As emulsifiers in baked goods and baking mixes, dairy product analogs, frozen desserts and mixes, and whipping milk products.
2. As texturizers in biscuit mixes.
3. As components of protective coating for fresh apples, bananas, pears, pineapples, avocados, plantains, limes, melons, papaya, peaches, and plums to retard ripening and spoilage.

The properties of SFE and olestra depend in part on the degree of substitution of sucrose with fatty acids, as shown in Table 4. SFE is used extensively in baked goods to improve the finished quality of frozen bread dough and sugar snap cookies. As a batter-aerating agent, SFE improves sponge and angel food cakes by increasing cake volume by 10–20%. Other functional users of SFE are shown in Table 5. These esters are seeing increased use in baked goods for several reasons:

1. They are easy to solubilize, can form good starch complexes, and can delay or control starch granule gelatinization. They prevent sticking to machinery and result in a softer crumb.
2. They are similar to the natural glycolipids found in wheat flour. Therefore, they promote expansion of gluten, stabilize aerated bubbles in batter, and make the manufacturing process easier.
3. They can inhibit crystal growth, such as ice crystals in frozen dough and sugar crystals in sweet baked goods.
4. They can increase loaf and cake volume and tenderness. They increase cookie spread factor and can affect crumb firmness and texture of sponge cakes.
Table 4  Some Properties of Sucrose Esters and Polyesters Based on Degree of Substitution

<table>
<thead>
<tr>
<th>Property</th>
<th>Degree of substitution</th>
<th>Approximate HLB value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrophilic</td>
<td>1–4</td>
<td>5–16</td>
</tr>
<tr>
<td>Lipophilic</td>
<td>5–8</td>
<td>1–3</td>
</tr>
<tr>
<td>Digestibility:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nondigestible</td>
<td>4–8</td>
<td>1–3</td>
</tr>
<tr>
<td>Digestible</td>
<td>1–3</td>
<td>5–16</td>
</tr>
<tr>
<td>Absorbability:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nonabsorbable</td>
<td>4–8</td>
<td>1–3</td>
</tr>
<tr>
<td>Absorbable</td>
<td>1–3</td>
<td>5–16</td>
</tr>
<tr>
<td>Emulsification:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>O/W emulsion</td>
<td>1–4</td>
<td>5–16</td>
</tr>
<tr>
<td>W/O emulsion</td>
<td>5–8</td>
<td>1–5</td>
</tr>
<tr>
<td>Antimicrobial</td>
<td>1–2</td>
<td>15–16</td>
</tr>
</tbody>
</table>

3. Antimicrobial Properties

The antimicrobial properties of SFE depend on the type and chain length of the fatty acid esterified to sucrose. Medium chain fatty acids such as lauric acid esterified to sucrose are better antimicrobial agents than sucrose esters of long chain fatty acids. Sucrose monoesters are more potent than the di-, tri-, and polyesters against gram-

Table 5  Some Functional Uses of Sucrose Fatty Acid Esters

<table>
<thead>
<tr>
<th>Product</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bread</td>
<td>Increase loaf volume and maintain softness</td>
</tr>
<tr>
<td>Noodles</td>
<td>Prevent sticking of mixed dough</td>
</tr>
<tr>
<td>Cake</td>
<td>Increase cake volume; shorten whipping time</td>
</tr>
<tr>
<td>Crackers, cookies</td>
<td>Stabilize emulsion; prevent sticking to machinery; increase volume</td>
</tr>
<tr>
<td>Ice cream</td>
<td>Improve overrun by promoting stable emulsion, thus preventing excessive cohesion of fat during freezing</td>
</tr>
<tr>
<td>Whipping cream</td>
<td>Prevent water separation</td>
</tr>
<tr>
<td>Margarine and W/O emulsions</td>
<td>Emulsification; prevent spattering</td>
</tr>
<tr>
<td>Shortening</td>
<td>Stabilize emulsion; increase water-holding capacity</td>
</tr>
<tr>
<td>O/W emulsions</td>
<td>Stabilize emulsions in a wide range of HLB values</td>
</tr>
<tr>
<td>Processed meat</td>
<td>Increase water-holding capacity of sausages; prevent separation of bolognas</td>
</tr>
<tr>
<td>Fruits</td>
<td>Coating to maintain freshness and extend shelf life</td>
</tr>
<tr>
<td>Drugs</td>
<td>Stabilize fat-soluble vitamins; lubricant; binder and filler</td>
</tr>
<tr>
<td>Cosmetics</td>
<td>Softness to skin; smoothness</td>
</tr>
<tr>
<td>Detergents</td>
<td>Cleaning agents for baby bottles; vegetables, and fruits</td>
</tr>
<tr>
<td>Antimicrobials</td>
<td>Prevent growth of microorganisms</td>
</tr>
</tbody>
</table>

*All used at no more than 10%.
positive bacteria. In general, SFEs exert more inhibitory action against gram-positive bacteria than against gram-negative bacteria. However, the polyesters are more active against gram-positive bacteria. The antimicrobial action of SFE is biostatic rather than biocidal in most cases. The antimicrobial properties of SFE received an excellent review from Marshall and Bullerman (38).

F. Esterified Propoxylated Glycerol

Esterified propoxylated glycerol (EPG) contains an oxypropylene group between the glycerol and fatty acids as follows:

$$EPIG = P(OH)_{a+c} \cdot (EPO)\cdot (FE)$$

where $P$ = polyol with two to eight primary OH groups (e.g., glucose), $C$ = zero to eight secondary and tertiary OH groups, $EPO$ = C3–C6 epoxide, $FE$ = acyl group of fatty acids C8–C24, $n$ = minimum epoxidation index average number ($\geq a$), $a + c$ = range of 3–8, and $b = \leq a + c$.

EPG is synthesized from glycerin and propylene oxide, which is subsequently esterified with fatty acids to yield oil-like product. The physical properties of the finished product, like natural triacylglycerols, depend on the type of fatty acid esterified. EPGs resemble triacylglycerols and can be used to replace fat in most food applications (39). Early results indicate that EPG is poorly hydrolyzed in animals fed the product for 30 days (9). It is anticipated that up to 3–4 more years of research is needed before ARCO Chemical Company will be able to petition the FDA for food additive status, with use in foods as a fat substitute contemplated. This compound may also have applications in the cosmetic and pharmaceutical industries.

G. Dialkyl Dihexadecylmalonate

Dialkyl dihexadecylmalonate (DDM) is synthesized from malonic acid, fatty acids, and hexadecane (40). It is a low-calorie fat substitute suitable for high temperature frying. DDM is absorbed, distributed, and eliminated through the liver. Frying test results indicate that there is no difference between potato and corn chips fried in a DDM–vegetable oil blend (60:40) and fried in conventional vegetable oil. Frito-Lay is expected to file a formal petition to the FDA for use of DDM as a fat substitute, upon completion of further testing and safety studies.

H. Jojoba Oil

Jojoba oil is composed of a mixture of linear esters of monounsaturated fatty acids and fatty alcohols containing 20–22 carbon atoms each (Fig. 14). It is liquid at 10°C and above. Nestec Limited (Switzerland) started research on the possible use of jojoba oil as a low-calorie food ingredient in 1979. Results indicate that jojoba oil is not fully hydrolyzed by the pancreatic lipases because of their structure. About

![Figure 14](Structure of jojoba oil.)
40% of the oil was absorbed by rats fed the oil (16). Safety and tolerance levels have been big issues. Indeed, rats consuming more than 16% of jojoba oil developed diarrhea and eventually died. Some levels of jojoba oil are stored in the liver. Possible use in control of obesity must be under careful medical supervision. No extreme adverse effects on liver, kidney, heart, or reproductive organ have been reported. Before jojoba could be marketed as a low-calorie oil, it would have to receive FDA approval (4).

I. Polycarboxylic Acid Esters and Ethers

Polycarboxylic acids containing two to four carboxylic acid groups esterified with saturated or unsaturated alcohols having straight or branched carbon chains consisting of 8–30 carbon atoms have been proposed as low-calorie edible oils (4,9,16,39,41). Because the ester groups are reversed (fatty alcohol esterified onto a polycarboxylic acid backbone) from the corresponding esters present in triacylglycerols, the polycarboxylic acid esters are not susceptible to complete hydrolysis by lipases. Hamm (16) also reported that trialkoxytricarballylate (TATCA, Fig. 15), trialkoxycitrate (TAC, Fig. 16), trialkoxyglyceryl ether (TGE, Fig. 17), and jojoba oil can serve as replacements for conventional edible fats and oils (Table 2). TATCA resembles triacylglycerol except that the glycerol backbone is replaced with tricarballylic acid and fatty acids with saturated and unsaturated alcohols.

TATCA is synthesized from tricarballylic acid and excess oleyl alcohol by solvent-free esterification at 135–150°C and a vacuum of 7 mm Hg. TAC is synthesized by essentially the same procedure as the TATCA except that molecular sieving
is not used to drive the reaction to the right (product formation). TATCA can replace vegetable oil in cooking and spreadable products such as margarine and mayonnaise. There are problems associated with the consumption of polycarboxylic acid esters. For example, in rat feeding experiments, anal leakage, weakness, depression, and death were observed when TATCA and jojoba oil were fed at moderate to high dose levels (1–3 g) (16). Weight gain data in rats indicated that both jojoba oil and TATCA are lower in calorie value than corn oil.

J. Polyglycerol Esters

The process for preparation and purification of polyglycerol and polyglycerol esters was described by Babayan (42,43). Esterification of glycerin with long chain fatty acids reduces absorption by 31 to 39% compared to corn oil, which is 98% absorbed. As the molecular weight of the polyglycerin portion increases, the hydrophilicity of the molecule also increases. But as the fatty acid chain length increases, hydrophilicity decreases. It was reported that male rats consuming 1 g/day of polyglycerol esters experienced weight gain comparable to that of rats fed lard. There were no abnormalities, except that diarrhea was observed in rats fed polyglycerol esters. Presently, polyglycerol esters are used as emulsifiers and dietetic acids such as in Weight Watchers ice cream. They can also be used in shortenings, margarines, bakery products, frozen desserts, ice cream, and confectioneries. The structure of a polyglycerol ester is shown in Figure 18.

K. Polysiloxane

Polysiloxane (PS) and phenylmethylpolysiloxane (a substituted polysiloxane) are organic derivatives of silica (SiO₂) with a linear polymeric structure (Fig. 19). The

![Figure 17](image17.png)

**Figure 17** Structure of trialkoxyglyceryl ether.

![Figure 18](image18.png)

**Figure 18** Structure of a polyglycerol ester.
polymeric molecules are inert, nontoxic, and varied in viscosity. Some have viscosity close to that of soybean oil. They are stable and not susceptible to oxidation as carbohydrate polyesters and vegetable oils. They are proposed as noncaloric, non-absorbable liquid oil substitutes (3). Studies in female obese Zucker rats resulted in weight reduction, and the animals did not compensate for the caloric dilution with PS by increasing their food intake (3).

IV. GENERAL PROPERTIES OF SYNTHETIC FAT SUBSTITUTES

Most of the physical properties studies have been conducted with sucrose fatty acid esters (SFE, DS < 4), which are digestible and absorbable. The surface-active properties of the mono- and diesters of sucrose were studied by Osipow et al. (30). These esters were found to be good emulsifying agents and detergents with low toxicity. SFEs may be useful in cosmetic, pharmaceutical, and food applications. They are soluble in warm water, ethanol, methanol, and acetone. Other properties reported include foaming, stability, wetting, softening point, percent acyl radical (determined by saponification of the sucrose esters), and surface and interfacial tension reduction. More information on the physical properties of sugar ester emulsifiers such as solubility, melting and decomposition temperatures, surface tension, solubilizing ability, foaming, and emulsification ability is available (34). The hydrophile–lipophile balance (HLB) of food-grade sucrose ester emulsifiers has been reported in two sources (34,44). The HLB of sucrose esters and sucrose ester–glyceride blends as emulsifiers was reported to be dependent on (a) degree of substitution or esterification (number of OH groups esterified with fatty acids), (b) alkyl chain length in the ester group, and (c) the presence of acyl double or triple bonds (i.e., degree of unsaturation). Sucrose esters are synergistic with sucrose polyesters of low HLB values in stabilizing O/W emulsions.

The physical properties of the more completely substituted carbohydrate and alkyl glycoside fatty acid polyesters (HLB ≥ 4) have recently been reported (10,12–14,45) and are summarized in Table 6. The color, consistency, density, specific gravity, and refractive indices of these polyesters approximate those of commercial vegetable and salad oils. However, the apparent viscosities of some of the carbohydrate polyesters were significantly greater than the apparent viscosities of salad oil. The HLB values of the carbohydrate polyesters were between HLB 2 and 6, suggesting that they are capable of promoting W/O emulsions and might be useful as emulsifiers in butter, margarine, low-fat spreads, caramel, chocolate, candy, and shortening. The
Table 6  Comparison of General Properties of Fat Replacers

<table>
<thead>
<tr>
<th>Property</th>
<th>Protein-based</th>
<th>Carbohydrate-based</th>
<th>Lipid-based</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physical form</td>
<td>Powder</td>
<td>Powder/liquid</td>
<td>Powder, solid, liquid</td>
</tr>
<tr>
<td>Taste</td>
<td>Bland</td>
<td>Bland</td>
<td>Bland</td>
</tr>
<tr>
<td>Caloric value, kcal/g</td>
<td>1–4</td>
<td>1–4.5</td>
<td>0–8.3</td>
</tr>
<tr>
<td>Melting point, °C</td>
<td>Variable</td>
<td>Variable</td>
<td>Range variable depending on fatty acid, unsaturation, and degree of substitution</td>
</tr>
<tr>
<td>Color</td>
<td>White</td>
<td>White</td>
<td>Yellow to golden yellow</td>
</tr>
<tr>
<td>Odor</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Soluble at 25°C in:</td>
<td>Water</td>
<td>Water, insoluble</td>
<td>Hexane, vegetable oils</td>
</tr>
<tr>
<td>Viscosity</td>
<td>Variable</td>
<td>Variable</td>
<td>Olestra more viscous than vegetable oils</td>
</tr>
<tr>
<td>Oxidative stability</td>
<td>Stable</td>
<td>Stable</td>
<td>Oil type of olestra not very stable; medium chain length triglycerides very stable</td>
</tr>
<tr>
<td>Stability to frying temperature</td>
<td>Not stable</td>
<td>Not stable</td>
<td>Most are stable</td>
</tr>
</tbody>
</table>

*Bland is a general term applicable to most but not all fat replacers.
*Variable melting point; depends on the specific compound. Actual values may be obtained from the supplier.

melting behavior of carbohydrate polyesters, such as sucrose and raffinose polyesters, has been reported (2,14,45). The melting points decreases with an increase in unsaturation of fatty acids. Carbohydrate or alkyl glycoside polyester with a desirable melting point range and other physical properties are prepared by blending fatty acid methyl esters of various degrees of saturation and unsaturation prior to synthesis of the saccharide polyester (14).

Lipid-based fat substitutes are subject to oxidation just like conventional triacylglycerols. The addition of the antioxidant TBHQ was reported to greatly improve the stability of liquid carbohydrate fatty acid polyester fat substitutes and vegetable oils. The degree of added stability was greater in the fat substitutes and in the refined, bleached, and deodorized (RBD) soybean oil than in the crude soybean oil (46). Part of the explanation was that RBD oils and fat substitutes made with them have lost some of the protective natural tocopherols, which must be restored to improve storage stability.

V. METABOLISM OF LIPID-BASED FAT SUBSTITUTES

Lipid-based fat substitutes, as exemplified by sucrose, trehalose, sorbitol, raffinose, stachyose, and sorbestrin, are not hydrolyzed by pancreatic lipase and consequently are not taken up by the intestinal mucosa (7,11,20,47). However, alkyl glycoside fatty acid polyesters may be partially hydrolyzed and absorbed (19). Mattson and Volpenhein (48) reported that as the number of ester groups increased from 4 to 8 in sucrose polyester, the rate of hydrolysis by lipase decreased. In other words, digestibility and absorbability of carbohydrate polyesters in rats and humans are
inversely related to the degree of substitution or esterification (20,48). Because of the molecular size of the fat substitutes and steric hindrance, the lipases cannot get to the substrate, and subsequently the formation of the enzyme–substrate [ES] complex required for lipase action on carbohydrate polyesters is prevented (11). Poor digestibility and absorbability of carbohydrate polyesters indicate that these substances supply few or zero calories; hence they are called low-calorie fat substitutes. Olestra is an FDA-approved, zero-calorie fat substitute. Because olestra is the most studied and the only synthetic lipid-based fat substitute approved, it is the focus of most of the discussions.

A. Effect of Olestra on Serum Lipids and Weight Reduction

An early report by Crouse and Grundy (49) indicated that sucrose polyester (olestra) interferes with the absorption of cholesterol because cholesterol is soluble in the fat substitute. According to Grundy et al. (50), nondiabetic patients on a calorie-restricted diet plus sucrose polyester (SPE) exhibited decreases in total cholesterol and low density lipoprotein (LDL) cholesterol of 20% and 26%, respectively. Reduction in serum cholesterol was at the expense of LDL-cholesterol. Diabetic patients with hypertriglyceridemia caloric restriction showed a marked reduction in plasma triacylglycerols with or without sucrose polyester consumption. Caloric restriction apparently reduced cholesterol by reducing cholesterol synthesis. Sucrose polyester has little effect on the concentration of HDL-cholesterol (47,51,52). LDL-cholesterol is associated with the development of atherosclerotic plaques, while HDL-cholesterol is associated with the prevention of atherosclerotic plaques. Patients with high cholesterol levels are advised to lower their serum cholesterol levels by losing weight and reducing both total fat intake (especially saturated fats) and cholesterol intake. Sucrose polyester may also be useful in weight reduction (11,19,47). Other carbohydrate polyesters can assist in the reduction of cholesterol levels (5,53–61). As cholesterol enters the digestive tract, it dissolves into the carbohydrate polyester oil phase and is excreted along with the carbohydrate polyester. Mellies et al. (53) reported that in obese hypercholesterolemic outpatients, SPE induced significant reductions in LDL-cholesterol beyond the effects of weight reduction. Glueck et al. (54) found that substitution of SPE for dietary fats in hypocaloric diets in obese women heterozygous for familial hypercholesterolemia resulted in 23% reduction of LDL-cholesterol and in weight loss. Mattson and Jandacek (58) reported that [4-\textsuperscript{14}C] cholesterol injected into rats that were subsequently fed SPE diets appeared in the feces of SPE-fed animals. Cholesterol absorption was reduced when 7 g of SPE was fed twice a day to 20 normocholesterolemic male inpatients in a double-blind crossover trial (56). It should be noted that Procter & Gamble did not pursue the approval of olestra as a cholesterol-reducing agent.

B. Effect of Olestra on Macronutrients and Fat-Soluble Vitamins

Olestra does not affect the availability of macronutrients such as carbohydrates, triacylglycerols, and proteins or that of micronutrients such as water-soluble vitamins, olestra consumption was, however, found to increase the intake of carbohydrate but without affecting total daily energy intake or usual patterns of hunger and fullness. SPEs may affect the availability of some fat-soluble vitamins such as vitamins A, D, E, and K, β-carotene, and lycopene (62), and cholesterol. In heavy snack eaters,
olestra can reduce vitamin A by 5% and vitamin E by about 3%. Addition of retinyl palmitate and tocopheryl acetate can offset the losses of vitamin A and E, respectively. The FDA requires that foods containing olestra be supplemented with fat-soluble vitamins to replace the amount normally found in vegetable oil and to compensate for olestra’s effect on vitamin absorption (Table 7).

Mellies et al. (52) reported earlier that plasma vitamin E was significantly reduced when 40 g/day of sucrose polyester was consumed by hypercholesterolemic outpatients compared to patients consuming a placebo. Crouse and Grundy (49) reported that vitamin E concentrations decreased by 24% and vitamin A was not affected. Fallat et al. (63) reported that both vitamins A and E were significantly reduced when SPE was added to patients diet at 50 g/wk of diet. In a later study, Mellies et al. (53) reported no decrease in vitamin A, but a 23% decrease in vitamin E. Daily consumption of 3 g of SPE led to significant reductions in plasma concentrations of β-carotene and lycopene in humans (62).

There is no clear consensus on the loss of vitamin A and carotenoids when consumed with olestra, and this may warrant further investigation. Vitamin D decreased in obese, hypercholesterolemic outpatients fed low-fat diets with or without SPE or conventional fat placebo (53). Miller et al. (64) supplemented both vitamins A and E in the diets of beagle dogs fed for 20 months with SPE at 10% and found no effect on the level of vitamins D and K, but concluded that with supplementation, vitamins A and E status remained sufficient in all groups. Another study suggested that vitamin K was not affected by consumption of SPE (52). Nutritional implications of fat substitute consumption and impact on fat intake were reported recently (9,65–68). Potential nutritional benefits of nondigestible lipid-based fat substitutes are shown in Table 8.

Carbohydrate polyesters in general, if taken in moderation, have the potential to help consumers lower their total fat intake, to reduce their saturated fat and cholesterol levels, to lose weight, and to achieve a healthier lifestyle. Again, they may benefit persons at high risk for coronary heart disease, colon cancer, and obesity (4,47,61).

C. Side Effects and Limitations

Anal leakage or oil loss resulted from consumption of large amounts of liquid olestra at the early stages of olestra development and studies. This can be prevented by

<table>
<thead>
<tr>
<th>Vitamin</th>
<th>Suggested supplementation level*</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>51 retinol eq/g olestra as retinyl palmitate or retinyl acetate = (170 IU/g olestra or 0.34 × RDA/10 g olestra)</td>
</tr>
<tr>
<td>D</td>
<td>12 IU vitamin D/g olestra = (0.3 × RDA/10 g olestra)</td>
</tr>
<tr>
<td>E</td>
<td>1.9 mg α-tocopherol eq/g olestra = (0.94 × RDA/10 g olestra)</td>
</tr>
<tr>
<td>K</td>
<td>8 µg vitamin K, g olestra = (1.0 × RDA/10 g olestra)</td>
</tr>
</tbody>
</table>

*IU, international unit; RDA, recommended daily allowance. The suggested levels are to compensate for amounts that are not absorbed from the diet because of olestra action. Source Adapted from Federal Register (51).

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Table 8  Some Nutritional Uses of Nondigestible Lipid-Based Fat Substitutes

<table>
<thead>
<tr>
<th>Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Replace saturated fat in the diet</td>
</tr>
<tr>
<td>Contribute reduced to zero calories to the diet</td>
</tr>
<tr>
<td>Reduce total fat intake in the diet</td>
</tr>
<tr>
<td>Reduce total cholesterol level</td>
</tr>
<tr>
<td>Reduce serum and plasma triacylglycerol levels</td>
</tr>
<tr>
<td>Reduce LDL-cholesterol level</td>
</tr>
<tr>
<td>Maintain ideal body weight or promote weight loss</td>
</tr>
<tr>
<td>Maintain HDL-cholesterol level</td>
</tr>
<tr>
<td>Reduce coronary heart disease risk factors</td>
</tr>
<tr>
<td>Reduce risk to certain kinds of cancer</td>
</tr>
</tbody>
</table>

addition of saturated fatty acids and their salts (59). It was suggested that oil loss can also be prevented by controlling the rheology (viscosity) and stiffness of carbohydrate polyesters by simply adjusting the fatty acid composition of the fats and oils used in their synthesis (11,19,51,61). Other gastrointestinal symptoms due to consumption of olestra are flatulence, soft stools, fecal urgency, diarrhea, and increased bowel movements (5,52,53). Indeed, it was recommended that soft stool can be prevented by using olestra with semisolid consistency or by increasing the viscosity of the olestra or product (51).

The most recently publicized problems with consuming olestra are its effects on serum carotenoid (antioxidant vitamin) levels, fat-soluble vitamins, gastrointestinal tract (oil loss, loose stools), and oral contraceptives or lipophilic drugs (Table 9). However, the role of carotenoids in health is still under investigation and not fully understood. Also not known is the effect of high levels of olestra consumption on consumers who may indulge in an effort to reduce or control their weight. To protect consumers, the FDA requires that products containing olestra bear labels (51) stating: “This Product Contains Olestra. Olestra may cause abdominal cramping and loose stools. Olestra inhibits the absorption of some vitamins and other nutrients.

Table 9  Limitations to Olestra Consumption

<table>
<thead>
<tr>
<th>Limitation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anal leakage or “oil loss” or fecal urgency</td>
</tr>
<tr>
<td>Decrease or loss of vitamins E and A from other foods</td>
</tr>
<tr>
<td>(FDA-recommended compensation, Ref. 51)</td>
</tr>
<tr>
<td>Soft stool</td>
</tr>
<tr>
<td>Diarrhea</td>
</tr>
<tr>
<td>Increased bowel movement</td>
</tr>
<tr>
<td>Loss of carotenoids (Ref. 62); 60% decrease in β-carotene</td>
</tr>
<tr>
<td>(supplementation under debate)</td>
</tr>
<tr>
<td>May affect drug absorption (coumarin, oral contraceptives—magnitude of effect small)</td>
</tr>
<tr>
<td>Inflammatory bowel disease</td>
</tr>
</tbody>
</table>

*FDA recommendations informing the consumer, through labeling, of some of these limitations, are given in Ref. 51.
Vitamins A, D, E, and K have been added.” Vitamins A, D, E, and K are required to be added as shown (Table 7) to protect consumers from depleting their essential vitamins (51). In other words, the added vitamins will compensate for the amounts that are not absorbed due to olestra’s interference with their absorption when olestra is eaten at the same time as foods containing the vitamins.

Olestra has the potential to benefit some segments of the population. From the available data, this substance is considered safe and nontoxic: because olestra is not digested or absorbed, no major component of it is available to produce unsafe or toxic effects (51). Research has shown that olestra is not genotoxic and does not affect reproduction or cause birth defects when consumed at certain levels. The long-term effects on humans of olestra use are open to debate. The published estimated daily intake (EDI) of olestra is 7 g/person for chronic consumption by the 90th percentile snack eaters and 20 g/person for short-term consumption (51).

Foods made with olestra compared with their commercially available counterparts taste the same, have the same flavor, texture, and appearance, and perhaps are less oily than the commercial product (51). Recent studies showed that olestra was not associated with increased incidence or severity of gastrointestinal symptoms (69,70).

VI. SAFETY AND REGULATORY UPDATES

For FDA approval of any new food additive, the petitioner must conduct toxicological, clinical, gastrointestinal, and nutritional testing to prove that the compound is safe and submit a food additive petition (FAP). Since the discovery of sucrose fatty acid polyester (olestra) in 1968 by Mattson and Volpenhein, extensive studies have been carried out in seven different species of animals, including long-term studies in monkeys and over 60 clinical trials involving over 8000 men, women, and children in several universities and medical research centers. The results show that olestra is safe: it is not absorbed or metabolized; it is nontoxic, nonmutagenic, and noncarcinogenic; and it does not affect reproduction or cause birth defects (48,59,62,71–75).

Olestra is biodegradable in sludge-amended soils and does not adversely affect the physical properties of the soil. It is not toxic to plants and animals. Gastrointestinal testing revealed that olestra does not affect gastrointestinal morphology, transit rate, bile acid physiology, bowel movement, pancreatic response, or intestinal microflora to any appreciable extent. It does not significantly affect the absorption or efficacy of orally dosed lipid-soluble drugs such as diazepam (tranquilizer), propranolol (cardiovascular agent), aspirin (analgesic), ethinyl estradiol (oral contraceptive), and noretindrone (oral contraceptive) (51).

The results reported by Miller et al. (64) indicated that sucrose polyester was not toxic when fed at 10% level to dogs in a 20-month study. No SPE was detected in the liver, heart, kidney, spleen, lymph nodes, and adipose tissues of 26 monkeys fed SPE at 0, 2, 4, or 6% on the diets for 2 months. Also, no SPE was detected in liver of these monkeys after 2 years of 9% SPE feeding. Daily consumption of 18 g of SPE did not affect the absorption or efficacy of the highly lipophilic oral contraceptives containing 300 μg of norgestrel and 30 μg of ethinyl estradiol (Lo/Oral-28) after 28 days (76). SPE is not genotoxic in the Salmonella/mammalian microsome test, the L5178Y thymidine kinate (TK +/−) mouse lymphoma assay, an unscheduled DNA synthesis array in primary rat hepatocytes, or an in vitro cyto-
VII. APPLICATIONS OF SUCROSE ESTERS

Sucrose fatty acid esters are sugar esters with a lower degree of substitution or esterification (DS 1–3). Sucrose fatty acid esters are FDA-approved and are currently marketed by Mitsubishi Chemical America, Inc. (White Plains, NY) and Crodesta, Inc. (New York). They are digestible and absorbable and can be used as emulsifiers and as lubricating, wetting or dispersion, solubilization, stabilization, anticaking, antimicrobial, or viscosity-reducing agents, in baking or as coatings for bananas, apples, pears, pineapples, and so on. Alkyl glycoside mono- and diesters can be used for similar applications. Table 5 shows some functional uses of sucrose fatty acid esters. The concentrations of SFE added to foods may range from 0.005% to 10%.

VIII. APPLICATIONS OF SYNTHETIC FAT SUBSTITUTES

Carbohydrate polyesters such as sucrose polyester has the potential to lower cholesterol levels in certain lipid disorders (50,52,53). Daily substitution of fat with 30 g of SPE will reduce caloric intake by 270 calories. Grundy et al. (50) reported that SPEs were tolerated by diabetic patients at a maximum of 90 g/day when total caloric intake was reduced to 1000 kcal/day. Thus, SPEs and other carbohydrate polyesters may be used for therapeutic weight reduction (47). They have the benefit of adding bulk to the diet without adding calories. Consumption of carbohydrate polyesters may benefit persons at high risk for coronary heart disease, colon cancer, and obesity (47,65). Table 8 lists some nutritional uses and applications of nondigestible fat substitutes.

Carbohydrate fatty acid polyesters are poor surfactants for O/W emulsions when used alone, but when blended with other emulsifiers or SFEs, they exhibit synergistic effects (2). Food made with SPE has 10–50% fewer calories than food made with regular oils and shortenings. For example, an order of french fries prepared with 100% SPE contains about 144 calories, 52.7% less than the same item with conventional oil. Thus, SPEs can be used for cooking, frying, and baking. They are useful in dairy products, spreads, dressings, margarines, sauces, confectioneries, and dessert manufacture.
<table>
<thead>
<tr>
<th>Item/property</th>
<th>Effect as petitioned</th>
<th>FDA ruling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stability</td>
<td>Stable as triacylglycerols; less polymer formation; higher free fatty acid formed</td>
<td>No difference in by-product formation during frying and baking</td>
</tr>
<tr>
<td>Absorption</td>
<td>Small amounts of penta- and lower esters are absorbed and metabolized to fatty acids and sucrose</td>
<td>Essentially not absorbed</td>
</tr>
<tr>
<td>Genetic toxicity</td>
<td>No evidence of genetic toxicity or mutagenicity from heated olestra</td>
<td>Not genotoxic</td>
</tr>
<tr>
<td>Teratogenicity</td>
<td>Not teratogenic or embryotoxic</td>
<td>No association to adenomas, pituitary, leukemia, basophilic foci with olestra treatment</td>
</tr>
<tr>
<td>Carcinogenicity</td>
<td>No adverse treatment-related effect</td>
<td>Not teratogenic</td>
</tr>
<tr>
<td>Drug absorption</td>
<td>No effect on lipophilic and nonlipophilic drug absorption</td>
<td>Magnitude of effect on drug absorption (including Coumadin, oral contraceptives) is not significant</td>
</tr>
<tr>
<td>Water-soluble nutrients</td>
<td>No adverse effect</td>
<td>No significant effect; no supplementation required</td>
</tr>
<tr>
<td>Fat-soluble vitamins</td>
<td>No significant decrease in vitamin A; dose-related significant decrease in vitamin E; small decrease in vitamin D; dose-response decrease in vitamin K&lt;sub&gt;1&lt;/sub&gt; 60% decrease in β-carotene</td>
<td>Not significant as analyzed Vitamin E decrease Serum vitamin D decrease Potential for vitamin K decrease on long-term olestra use Carotenoids’ effect not conclusive; supplementation for vitamins A, D, E, K recommended</td>
</tr>
<tr>
<td>Gastrointestinal tract</td>
<td>GI symptoms not clinically significant</td>
<td>Dose–response for diarrhea/loose stools and fecal urgency; recommend informing consumer through labeling</td>
</tr>
<tr>
<td></td>
<td>Oil loss if large liquid olestra is consumed</td>
<td>Low stiffness leads to oil loss, not a health hazard</td>
</tr>
<tr>
<td></td>
<td>Increased bowel movement, soft stools</td>
<td>Symptoms not the same as in high-fiber diets</td>
</tr>
<tr>
<td></td>
<td>Inflammatory bowel disease affected little</td>
<td>Not highly exacerbated at 20 g/day</td>
</tr>
<tr>
<td></td>
<td>No significant health effects on young children</td>
<td>Data from adults can be extrapolated to children</td>
</tr>
<tr>
<td>Intestinal microflora</td>
<td>Olestra is not metabolized by GI microflora; microbial counts, short chain fatty acid, breath hydrogen not greatly affected</td>
<td>No metabolism by microflora; no differences as observed with dietary changes</td>
</tr>
<tr>
<td>Bile acid metabolism</td>
<td>No effect on bile acid synthesis or excretion</td>
<td>No major changes in bile acid metabolism</td>
</tr>
</tbody>
</table>
IX. PERSPECTIVES

With the approval of olestra, it is expected that studies by other competitors will be accelerated and alternative compounds will be developed to challenge the olestra market. Other approval requests for uses of olestra are expected to be made if the test market results for olestra-made snacks is favorable. However, if more problems arise from the test marketing, more opposition to requested approvals is expected. Approval for a new food additive is not easy: the additive petition for olestra/Olean contained over 150,000 pages that included safety data on more than 100 laboratory studies on seven animal species and more than 60 clinical trials involving over 8000 men, women, and children. Therefore, any new food additive or new use of an approved ingredient must undergo the same FDA petition and approval process, and studies must include toxicologic, preclinical, and clinical data showing that the substance is safe and efficacious.

It appears that consumers are willing to try low-fat or reduced-calorie foods (6). The market for fat substitutes is likely to increase. The labeling requirement for fat is likely to be revised to include nondigestible fat substitutes and reduced-calorie structured lipids. Foods containing fat substitutes will be so labeled to inform the consumer of potential benefits and health implications. Obviously, more research is needed in this area of consumer interest. The FDA and the government must be willing to support more research through independent investigators in the interest of the consumer. The need for more studies on the long-term effect of fat substitutes on the immune system, tolerance levels/dosage, essential fatty acid availability, degradation in the soil, physical properties, applications, stability, and analysis in food matrices should be emphasized.

Reduced-calorie fats and fat substitutes are likely to offer the public new strategies and more choices for reducing fat consumption. The specialty or structured lipids may compete with the zero-calorie fats or may come to be preferred because they can be designed for optimal nutrition, health, and disease prevention or to contain fewer calories than conventional fats and oils (see Chaps. 28 and 30). A partial replacement and systems approach, rather than total replacement of dietary fats with olestra-type materials, and consumption in moderation, should be encouraged. Total elimination of fat in the diet is not nutritionally prudent. A systems approach to formulating reduced-, low-, or no-calorie foods seems appropriate at this time, barring the absence of a “magic bullet” or an “ideal” fat substitute.

REFERENCES


I. FATS AVAILABLE FOR FOOD APPLICATIONS

A. Introduction

The annual production of oils and fats now exceeds 113 million tonnes and is on a rising curve. Table 1 contains data for the four major vegetable oils, some minor vegetable oils, and the animal fats. These are average annual oil production figures for a series of 5-year periods including some predictions for the future and show the increasing dominance of vegetable oils and particularly of soybean oil, palm oil, rapeseed/canola oil, and sunflower oil. The information is presented on a 5-year basis to avoid unusual harvests in any particular year. This book is devoted to the 80% of total lipids that are part of the human diet. An additional 6% is fed to animals, and the remaining 14% is the basis of the oleochemical industry, which produces mainly a wide range of surface-active compounds some of which are important in the food industry.

*Oil World* (1) provides regular statistics on 17 oils and fats, a list that excludes cocoa butter (Sec. VI), rice bran oil, and many specialty oils. The 17 oils vary in their fatty acid and triacylglycerol composition and have differing physical, chemical, and nutritional properties. The food technologist has to work with these resources to produce a wide range of products with optimized properties. Similar products can frequently be obtained with more than one selection of oils and fats, as is apparent in the fact that similar materials available in different regions of the world are made from different blends of natural or processed oils. But what happens when the natural products do not provide the necessary range of properties? Lipid scientists and technologists have devised several procedures to extend the range of natural fats. These are listed in Table 2. Some are discussed in greater detail in other chapters in this book, and they have been described elsewhere by the present author (2,3).
Table 1  Average Annual Production (million metric tonnes) of 17 Commodity Oils Over Selected 5-Year Periods Along with Future Projections

<table>
<thead>
<tr>
<th></th>
<th>1976/80</th>
<th>86/90</th>
<th>96/00</th>
<th>06/10</th>
<th>16/20</th>
</tr>
</thead>
<tbody>
<tr>
<td>17 oils and fats</td>
<td>52.6</td>
<td>75.7</td>
<td>103.4</td>
<td>139.4</td>
<td>175.8</td>
</tr>
<tr>
<td>Soybean</td>
<td>11.2</td>
<td>15.3</td>
<td>22.8</td>
<td>30.4</td>
<td>37.1</td>
</tr>
<tr>
<td>Palm</td>
<td>3.7</td>
<td>9.2</td>
<td>17.9</td>
<td>29.2</td>
<td>40.8</td>
</tr>
<tr>
<td>Rapeseed</td>
<td>3.0</td>
<td>7.5</td>
<td>12.6</td>
<td>17.5</td>
<td>22.2</td>
</tr>
<tr>
<td>Sunflower</td>
<td>4.2</td>
<td>7.2</td>
<td>9.1</td>
<td>12.5</td>
<td>16.6</td>
</tr>
<tr>
<td>Lauric oils (2)</td>
<td>3.3</td>
<td>4.3</td>
<td>5.4</td>
<td>7.5</td>
<td>9.6</td>
</tr>
<tr>
<td>Other veg oils (7)</td>
<td>10.0</td>
<td>12.3</td>
<td>14.9</td>
<td>18.0</td>
<td>21.6</td>
</tr>
<tr>
<td>Animal fats (4)</td>
<td>17.2</td>
<td>19.9</td>
<td>20.7</td>
<td>24.3</td>
<td>27.9</td>
</tr>
</tbody>
</table>

*These figures are taken from *Oil World 2020* (1). Averages of 5-year periods are taken to avoid unusually high or low figures such as occur in some years.
Lauric oils: palm kernel and coconut; other vegetable oils: cottonseed, groundnut, sesame, corn, olive, linseed, and castor; animal fats: butter, lard, tallow, and fish oil.

B. Fatty Acid Composition of Major Oils

Palmitic (16:0), oleic (18:1), and linoleic acids (18:2) are the dominant fatty acids in vegetable oils, and many oils and fats differ only in the relative proportion of these three acids. However, other fatty acids do exist and may be important. Stearic acid (18:0) is a minor component in virtually all oils; linolenic acid (18:3) becomes important in some; and the so-called lauric oils are rich in lauric (12:0) and myristic acids (14:0) (Table 3). Of the processes listed in Table 2 it is important to note that blending and fractionation lead to changes in the proportions of fatty acids. Interesterification does not change fatty acid composition but modifies the ways in which these are incorporated into triacylglycerols. However, (partial) hydrogenation results in the formation of novel fatty acids with structures different from those present in

Table 2  Methods Employed to Extend the Usefulness and Improve the Properties of Natural Oils

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blending</td>
<td>Mixing of two or more oils</td>
</tr>
<tr>
<td>Fractionation</td>
<td>Separation of oils into two or more fractions</td>
</tr>
<tr>
<td>Partial hydrogenation</td>
<td>Saturation of some double bonds accompanied by double-bond isomerization</td>
</tr>
<tr>
<td>Interesterification with chemical or enzymatic catalyst</td>
<td>Reorganization of fatty acids among triacylglycerol molecules</td>
</tr>
<tr>
<td>Domestication of wild crops</td>
<td>Conversion of wild crops to crops that can be cultivated commercially</td>
</tr>
<tr>
<td>Seed breeding by traditional methods</td>
<td>Interspecies crossing using irradiation or mutagenesis if necessary</td>
</tr>
<tr>
<td>Seed breeding by genetic modification</td>
<td>Crossing between species</td>
</tr>
<tr>
<td>Lipids from microorganisms</td>
<td>Production under fermentation conditions</td>
</tr>
</tbody>
</table>
Table 3  Typical Fatty Acid Composition (% wt) of Four Major Vegetable Oils, Lauric Oils, Animal Fats, and Hydrogenated Soybean Oil

<table>
<thead>
<tr>
<th></th>
<th>16:0</th>
<th>18:0</th>
<th>18:1</th>
<th>18:2</th>
<th>Other acids</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soybean</td>
<td>11</td>
<td>4</td>
<td>22</td>
<td>53</td>
<td>18:3 8%</td>
</tr>
<tr>
<td>Soybean (hydrog.)(^a)</td>
<td>11</td>
<td>8–12</td>
<td>63–72</td>
<td>9–13</td>
<td>18:3 &lt;0.3%</td>
</tr>
<tr>
<td>Palm</td>
<td>44</td>
<td>4</td>
<td>40</td>
<td>10</td>
<td></td>
</tr>
<tr>
<td>Rapeseed</td>
<td>4</td>
<td>2</td>
<td>56</td>
<td>26</td>
<td>18:3 10%</td>
</tr>
<tr>
<td>Sunflower (high-lin.)</td>
<td>6</td>
<td>5</td>
<td>20</td>
<td>69</td>
<td></td>
</tr>
<tr>
<td>NuSun(^b)</td>
<td>4–5</td>
<td>3–4</td>
<td>50–75</td>
<td>20–30</td>
<td></td>
</tr>
<tr>
<td>High-oleic</td>
<td>4</td>
<td>5</td>
<td>81</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>Olive</td>
<td>10</td>
<td>2</td>
<td>78</td>
<td>7</td>
<td>8:0 8%, 10:0 7%</td>
</tr>
<tr>
<td>Coconut</td>
<td>9</td>
<td>2</td>
<td>7</td>
<td>2</td>
<td>12:0 48%, 14:0 16%</td>
</tr>
<tr>
<td>Palm kernel</td>
<td>9</td>
<td>3</td>
<td>15</td>
<td>2</td>
<td>8:0 4%, 10:0 4%</td>
</tr>
<tr>
<td>Butter</td>
<td>26</td>
<td>11</td>
<td>28</td>
<td>2</td>
<td>12:0 45%, 14:0 18%</td>
</tr>
<tr>
<td>Lard</td>
<td>27</td>
<td>11</td>
<td>41</td>
<td>11</td>
<td>4:0–12:0 13%, 14:0 12%</td>
</tr>
<tr>
<td>Beef tallow</td>
<td>27</td>
<td>7</td>
<td>48</td>
<td>2</td>
<td>4:0 2%, 16:1 4%</td>
</tr>
</tbody>
</table>

\(^a\)Hydrogenated soybean oil of iodine value ~80 (normally about 133). 18:1 includes 27–40% of trans isomers.

\(^b\)NuSun is a sunflower oil with the levels of oleic acid shown in the table.

the native oils. New acids are formed with double bonds of changed configuration and changed position in the carbon chain.

C. Availability of Oils and Fats in Different Regions of the World

In the days before extensive world trade in perishable goods, foods were based on local availability and food traditions became established. These are still apparent throughout the world.

- Communities living near the seashore or a lakeside often consume more fish in their diet than other communities not so favorably placed with respect to these healthy foods.
- Countries and regions that produce large quantities of a particular oil generally consume a large amount of that material. This is very apparent in certain developing countries but it is also true of the developed world. It has been reported that 82% of the fat consumed in the United States is soybean oil (before or after some degree of hydrogenation) and that no other oil exceeds 3% (4). Some relevant data are presented in Table 4 but these need careful interpretation. “Disappearance” for a country or region represents locally produced oil plus imported oil minus exported oil with a small adjustment for changes in stock levels at the beginning and end of the reporting season. The figures include food and nonfood uses and make no allowance for waste of commodities outside the 17 listed in the reports. As already indicated, on a worldwide basis about 20% of all oils and fats are used for nonfood purposes. This proportion will be lower in many de-
Table 4  Annual Average Disappearance of Oils and Fats in 1996/00 in Selected Countries in Percentage Terms

<table>
<thead>
<tr>
<th>Country</th>
<th>sb</th>
<th>palm</th>
<th>rape</th>
<th>sun</th>
<th>pk</th>
<th>co</th>
<th>tl</th>
<th>lard</th>
<th>bt</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Canada</td>
<td>19.8</td>
<td>0.7</td>
<td>46.5</td>
<td>2.1</td>
<td>0.7</td>
<td>1.0</td>
<td>6.7</td>
<td>6.8</td>
<td>4.8</td>
<td>10.9</td>
</tr>
<tr>
<td>USA</td>
<td>51.9</td>
<td>1.0</td>
<td>4.8</td>
<td>0.6</td>
<td>0.6</td>
<td>1.4</td>
<td>4.0</td>
<td>19.1</td>
<td>3.2</td>
<td>10.9</td>
</tr>
<tr>
<td>Arg.</td>
<td>8.4</td>
<td>0</td>
<td>56.3</td>
<td>0.5</td>
<td>0.5</td>
<td>16.5</td>
<td>2.9</td>
<td>5.4</td>
<td>9.5</td>
<td></td>
</tr>
<tr>
<td>Brazil</td>
<td>70.3</td>
<td>2.6</td>
<td>1.9</td>
<td>0</td>
<td>1.9</td>
<td>1.4</td>
<td>0</td>
<td>5.5</td>
<td>2.0</td>
<td>7.3</td>
</tr>
<tr>
<td>Indon.</td>
<td>0.6</td>
<td>88.6</td>
<td>0</td>
<td>0</td>
<td>3.0</td>
<td>0</td>
<td>6.8</td>
<td>0</td>
<td>0</td>
<td>1.0</td>
</tr>
<tr>
<td>Malay.</td>
<td>2.0</td>
<td>60.2</td>
<td>0.3</td>
<td>0.3</td>
<td>34.0</td>
<td>1.7</td>
<td>0.1</td>
<td>0.1</td>
<td>0.4</td>
<td>1.0</td>
</tr>
<tr>
<td>Phil.</td>
<td>11.1</td>
<td>16.0</td>
<td>0.2</td>
<td>0.2</td>
<td>0.9</td>
<td>9.1</td>
<td>3.3</td>
<td>10.2</td>
<td>1.9</td>
<td>1.1</td>
</tr>
<tr>
<td>EU-15</td>
<td>11.6</td>
<td>11.4</td>
<td>14.7</td>
<td>12.6</td>
<td>2.3</td>
<td>3.8</td>
<td>8.7</td>
<td>8.2</td>
<td>9.3</td>
<td>17.4</td>
</tr>
<tr>
<td>Japan</td>
<td>25.4</td>
<td>13.8</td>
<td>32.6</td>
<td>0.5</td>
<td>2.1</td>
<td>1.9</td>
<td>5.0</td>
<td>2.6</td>
<td>2.8</td>
<td>13.3</td>
</tr>
<tr>
<td>India</td>
<td>10.8</td>
<td>16.6</td>
<td>21.0</td>
<td>6.9</td>
<td>0</td>
<td>4.3</td>
<td>1.5</td>
<td>0.2</td>
<td>12.8</td>
<td>25.9</td>
</tr>
<tr>
<td>China</td>
<td>21.6</td>
<td>10.1</td>
<td>24.0</td>
<td>1.2</td>
<td>0</td>
<td>0.3</td>
<td>4.8</td>
<td>16.2</td>
<td>0.5</td>
<td>21.3</td>
</tr>
</tbody>
</table>

Oils and fats: sb, soybean; rap, rapeseed/canola; sun, sunflower; pk, palm kernel; co, coconut; tl, tallow; bt, butter. Countries: Argentina, Indonesia, Malaysia, Philippines.

• The figures in Table 4 show that there are several countries in which an oil and fat grown locally is also the major fat, consumed at a level close to or higher than 50% of total disappearance, e.g., Canada canola oil 47%, USA soybean oil 52%, Philippines coconut oil 55%, Argentina sunflower oil 56%, Malaysia palm oil 60%, Brazil soybean oil 70%, and Indonesia palm oil 89%. In most cases the real figures will be higher than those cited above when allowance has been made for the use of lauric oils and tallow in the oleochemical industry (see above for the position in the USA). The situation is very different in countries that import large amounts of seed for local crushing or import oils and fats. This is apparent in the figures in Table 4 for EU-15, Japan, China, and India. EU-15 is an association of states covering a wide geographic region, and the figures in Table 5 reveal that there

Table 5  Disappearance of Oils and Fats During 1999 in Selected EU-15 Countries in Percentage Terms

<table>
<thead>
<tr>
<th>Country</th>
<th>sb</th>
<th>palm</th>
<th>rape</th>
<th>sun</th>
<th>pk</th>
<th>co</th>
<th>olive</th>
<th>tl</th>
<th>lard</th>
<th>bt</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>UK</td>
<td>7.8</td>
<td>22.3</td>
<td>33.2</td>
<td>5.2</td>
<td>4.0</td>
<td>2.1</td>
<td>1.4</td>
<td>8.0</td>
<td>2.5</td>
<td>9.0</td>
<td>4.5</td>
</tr>
<tr>
<td>Ger.</td>
<td>10.3</td>
<td>12.4</td>
<td>27.9</td>
<td>7.0</td>
<td>6.1</td>
<td>7.1</td>
<td>0.9</td>
<td>0.9</td>
<td>11.0</td>
<td>13.2</td>
<td>3.2</td>
</tr>
<tr>
<td>Fr.</td>
<td>2.9</td>
<td>5.3</td>
<td>20.7</td>
<td>23.0</td>
<td>0.7</td>
<td>2.7</td>
<td>3.9</td>
<td>3.8</td>
<td>8.1</td>
<td>21.7</td>
<td>7.2</td>
</tr>
<tr>
<td>Italy</td>
<td>13.7</td>
<td>8.4</td>
<td>3.9</td>
<td>10.1</td>
<td>1.1</td>
<td>2.1</td>
<td>31.6</td>
<td>6.0</td>
<td>8.6</td>
<td>4.9</td>
<td>9.6</td>
</tr>
<tr>
<td>Spain</td>
<td>11.6</td>
<td>5.5</td>
<td>1.9</td>
<td>25.5</td>
<td>1.3</td>
<td>1.3</td>
<td>26.4</td>
<td>8.5</td>
<td>13.2</td>
<td>0.9</td>
<td>3.9</td>
</tr>
</tbody>
</table>

Abbreviations: see Table 4. Countries: United Kingdom, Germany, France.
are significant differences in oil and fat disappearance between individual countries and particularly between those in the north and the south. Rapeseed oil, sunflower oil, and olive oil are strongly used in the European countries in which they are produced. Butter is still a significant fat in Europe except in the Mediterranean countries producing olive oil. The choice open to European food producers and consumers is apparent in the fact that no figure in Table 5 exceeds 35%.

With the free movement of oilseeds and of oils, and fats, trading nations have a wide choice among these commodities. With the development of high-quality refined oils many of them become interchangeable though the attention must be paid to fatty acid and triacylglycerol composition; physical, nutrition, and chemical properties; price; and customer concerns. Food producers monitor their supplies to make best use of the cheapest oils, and when customers express concerns, such as those over oils from genetically modified sources (mainly soybean oil) or oils containing trans acids, producers switch to recipes that do not need such materials. The message here is that while the major producing countries make extensive use of local products, there are regions of the world in which flexibility of supply is possible and is cherished by producers and consumers alike.

II. FRYING OILS AND FATS

The use of oils and fats as a frying medium in both shallow and in deep frying mode is an important component in the overall picture of food applications. It has recently been reported that 20 million tonnes of oils and fats is used in this way. This represents a major share of the 90 million tonnes used for dietary purposes. Of course, it must be remembered that while some of the frying oil is consumed along with the fried food, much is thrown away (shallow pan frying) or ultimately finds other uses as spent frying oil. The importance of frying is reflected by the fact that this topic has a chapter to itself, and this matter will not be pursued here except to provide some general references that may have been overlooked in Chapter 7 (5–9).

III. SPREADS: BUTTER, GHEE, MARGARINE, VANASPATI

A. Butter

Butter from cow’s milk fat has been used primarily as a spread, but also for baking and frying, for many centuries. With the development of good-quality margarine and other spreads butter has become less popular. The disadvantages associated with butter are its relatively high price, its poor spreadability (especially from the refrigerator), and its low health profile resulting from its high fat content, its high level of saturated acids and of cholesterol, and the presence of trans unsaturated fatty acids. Its advantages are its “wholly natural” profile and its superb flavor. The perceived superiority of butter is reflected in the fact that spread manufacturers seek to have products with the flavor and appearance of butter and by their use of names like “I Can’t Believe It’s Not Butter” for one brand of spread. Although the name butter is jealously guarded and it is not permissible to take anything away or to add anything to a product that is to be called butter, nevertheless ways of overcoming
the disadvantages are being reported. References 10–13 provide general information on butter.

Butter is a water-in-oil emulsion consisting of fat (80–82%) and an aqueous phase (18–20%). The legal limit for water is 16% and the aqueous phase also contains salt and milk-solids-not-fat. It is made from cow’s milk (3–4% fat), which is converted first to cream (30–45% fat) by centrifuging and then to butter by churning and kneading. During churning there is a phase inversion from an oil-in-water to a water-in-oil emulsion. Details of annual production and disappearance are given in Table 6. Production of butter, which peaked in the 1980s (6.0–6.4 MMT) has now fallen below 6 MMT but is predicted to rise steadily to 7.8 MMT in the next 20 years. The biggest consumers are in the Indian subcontinent (India and Pakistan) and in EU-15 within which France has a consumption equivalent to more than 20% of total oil and fat disappearance for that country (Table 5).

The composition of milk fat changes between summer and winter according to changing dietary intake (Table 7). The fat is mainly tricacylglycerols (97–98%) along with some free acids, monoacylglycerols, and diacylglycerols. Also present are cholesterol (0.2–0.4%), phospholipids (0.2–1.0%), and traces of carotenoids, squalene, and vitamins A and D.

Following several thorough examinations, cow’s milk fat is now known to contain more than 500 different fatty acids. Most of these are present only at exceedingly low levels but some of these, such as the lactones which provide important flavor notes, are important. Among the many fatty acids are the following (see also Table 7):

- Saturated acids in the range 4:0 to 18:0 including some odd-chain members.
- Low levels of iso-, anteiso-, and other branched chain acids.
- A significant level of monoene acids (28–31%) that is mainly oleic but includes other isomers, among which are some with trans configuration.
- trans acids, produced by biohydrogenation of dietary lipids, are significant components of all ruminant milk fats (about 4–8%). They are mainly C_{16} and C_{18} monoene acids of which vaccenic acid (11t-18:1) is the major component.
- Very low levels of polyene fatty acids and even those cited as linoleic or linolenic are not entirely the all-cis isomers.
- Trace amounts of oxo (keto) and hydroxy acids and lactones of which the latter are important flavor components.

The fatty acid composition of milk depends on the diet of the cow, so that in many countries there is a difference in composition between winter (fed indoors) and summer (pasture fed), and this can be further modified by controlling the diet. An exciting development in lipid science in recent years has been the recognition of the importance of certain octadecadienoic acids with conjugated unsaturation (conjugated linoleic acid, CLA), which are produced by ruminants and appear in low but significant levels in the milk and meat of these animals. This topic is important enough to merit a chapter of its own in this volume (Chapter 21).

The level of butyric acid at around 4% by weight may be considered insignificant, but it should be recognized that this is equivalent to about 8.5% mol. Since this acid is likely to occur only once in any triacylglycerol molecule (at the sn-3 position), a quarter of all the triacylglycerol molecules in butter contain butyric acid.
Table 6  Production and Disappearance (Million Metric Tonnes) of Butter on a Fat Basis for Selected Countries, 1995–1999

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>World</td>
<td>5.68</td>
<td>5.65</td>
<td>5.69</td>
<td>5.74</td>
<td>5.83</td>
<td>5.73</td>
<td>5.66</td>
<td>5.72</td>
<td>5.75</td>
<td>5.81</td>
</tr>
<tr>
<td>EU-15</td>
<td>1.56</td>
<td>1.60</td>
<td>1.56</td>
<td>1.58</td>
<td>1.59</td>
<td>1.42</td>
<td>1.52</td>
<td>1.47</td>
<td>1.51</td>
<td>1.53</td>
</tr>
<tr>
<td>USSR</td>
<td>0.69</td>
<td>0.54</td>
<td>0.48</td>
<td>0.46</td>
<td>0.45</td>
<td>0.83</td>
<td>0.64</td>
<td>0.64</td>
<td>0.56</td>
<td>0.51</td>
</tr>
<tr>
<td>E. Europe</td>
<td>0.21</td>
<td>0.21</td>
<td>0.20</td>
<td>0.21</td>
<td>0.20</td>
<td>0.19</td>
<td>0.17</td>
<td>0.19</td>
<td>0.19</td>
<td>0.18</td>
</tr>
<tr>
<td>USA</td>
<td>0.47</td>
<td>0.43</td>
<td>0.43</td>
<td>0.39</td>
<td>0.43</td>
<td>0.45</td>
<td>0.42</td>
<td>0.42</td>
<td>0.40</td>
<td>0.44</td>
</tr>
<tr>
<td>India</td>
<td>1.15</td>
<td>1.20</td>
<td>1.24</td>
<td>1.28</td>
<td>1.30</td>
<td>1.15</td>
<td>1.20</td>
<td>1.25</td>
<td>1.28</td>
<td>1.31</td>
</tr>
<tr>
<td>Pakistan</td>
<td>0.37</td>
<td>0.39</td>
<td>0.41</td>
<td>0.42</td>
<td>0.43</td>
<td>0.37</td>
<td>0.39</td>
<td>0.42</td>
<td>0.42</td>
<td>0.43</td>
</tr>
<tr>
<td>New Zealand</td>
<td>0.23</td>
<td>0.25</td>
<td>0.29</td>
<td>0.29</td>
<td>0.29</td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
</tr>
</tbody>
</table>

Most of the saturated acids (C₄–C₁₄ and about one-half of the C₁₆) are produced by de novo synthesis in the mammary gland while the rest of the C₁₆ and the saturated and unsaturated C₁₈ acids are derived from dietary sources or by mobilization of body fat reserves during early lactation. It follows that only this half of milk fat fatty acids is subject to change by modification of dietary intake. Furthermore, since in ruminants (free) unsaturated acids are subject to biohydrogenation in the rumen, it is necessary to protect such acids during their passage through the rumen if these are to be incorporated unchanged in the milk fat. This was first achieved through coating the oil/fat (soybean, linseed, rapeseed/canola), but two other methods are now more commonly employed. In the first, calcium salts are used as lipid source. These remain as (unreactive) salts in the rumen but are converted to acids in the more acidic conditions of the abomasum and enter the duodenum as fatty acids available for digestion. Alternatively, the lipid is hardened to the point where it remains solid in the rumen but melts in the abomasum. The resulting changes in the milk fat may seem small in terms of fatty acid composition, but are slightly greater in their effect on triacylglycerol composition and may be sufficient to allow the butter to spread directly from the refrigerator. It is important that the supplement contain appropriate proportions of n-9, n-6, and n-3 unsaturated acids and that it is over 75% protected from metabolism in the rumen (14,15).

In times of oversupply there is an interest in extending the range of applications of milk fat by fractionation. However, the triacylglycerol composition of milk fat is so complex (no individual triacylglycerol exceeds 5%) that differences between crystallized fractions are not so marked as with simpler vegetable oils, such as palm oil.

Table 7  Major Fatty Acids (% wt) in Cow’s Milk Fat

<table>
<thead>
<tr>
<th>Month</th>
<th>4:0</th>
<th>6:0</th>
<th>8:0</th>
<th>10:0</th>
<th>12:0</th>
<th>14:0</th>
<th>16:0</th>
<th>18:0</th>
<th>20:0</th>
<th>18:1</th>
<th>18:2</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>June</td>
<td>4.2</td>
<td>2.5</td>
<td>2.3</td>
<td>2.2</td>
<td>2.4</td>
<td>9.0</td>
<td>22.1</td>
<td>14.3</td>
<td>2.6</td>
<td>30.4</td>
<td>1.2</td>
<td>6.8</td>
</tr>
<tr>
<td>December</td>
<td>3.5</td>
<td>2.2</td>
<td>1.1</td>
<td>2.6</td>
<td>2.8</td>
<td>10.6</td>
<td>26.0</td>
<td>11.6</td>
<td>2.3</td>
<td>24.8</td>
<td>2.8</td>
<td>9.7</td>
</tr>
<tr>
<td>Average</td>
<td>3.6</td>
<td>2.2</td>
<td>1.2</td>
<td>2.8</td>
<td>2.8</td>
<td>10.1</td>
<td>25.0</td>
<td>12.1</td>
<td>2.1</td>
<td>27.1</td>
<td>2.4</td>
<td>8.6</td>
</tr>
</tbody>
</table>
Nevertheless, useful separations have been achieved producing fractions that are harder and softer than the original milk fat. The lower melting (softer) fractions are employed to make spreadable butter and the harder fractions find pastry applications. Anhydrous milk fat itself is used to make cakes. Mixed with the oleic fraction it is used in cookies, biscuits, and butter cream; mixed with the stearic fraction it is used in fermented pastries and puff pastry; and the oleic fraction on its own is used in ice cream cones, waffles, butter sponges, and in chocolate for ice cream bars (16,17).

In Europe, butters with reduced fat levels (and therefore reduced caloric values) are designated as butter (80–90% fat), three-quarters fat butter (60–62% fat), half fat butter (39–41% fat), and dairy fat spreads (other fat levels). In the United States, “light butter” must contain less than half of the normal level of fat and “reduced butter” less than one-quarter of the normal level.

Products are available in many countries that are blends of butter and vegetable oil—generally soybean oil. These cannot be called butter but are given an appropriate name that the consumer comes to think of as “spreadable butter.” Spreadable butters developed in New Zealand are made by fractionation of butter followed by recombination of appropriate fractions.

B. Ghee

In India, milk fat is consumed partly as butter but also as ghee, though the latter is declining and is now probably below one-quarter of the combined total. Nevertheless, demand in other countries is growing, probably reflecting the migration of Indians. Ghee is a concentrate of butterfat with more than 99% milk fat and less than 0.2% moisture. It has a shelf life of 6–8 months even at ambient tropical temperatures. Butter or cream is converted to ghee by controlled heating to reduce the water content to below 0.2%. In other procedures the aqueous fraction is allowed to separate and some of it is run off before residual moisture is removed by heating. Ghee has a cooked caramelized flavor varying slightly with the method of preparation (10,18). The vegetable oil–based alternative to ghee is called vanaspati (Sec. III.D.)

C. Margarine

Margarine has been produced for more than 100 years. During the 1860s, large sections of the European population migrated from country to town and changed from rural to urban occupations. At the same time, there was a rapid increase in population in Europe and a general recession in agriculture leading to a shortage of butter, especially for the growing urban population. As a consequence, the price rose beyond the reach of many poor people. So bad was the situation in France that the government offered a prize for the best proposal for a butter substitute that would be cheaper and would also keep better.

The prize was won by the French chemist, Hippolyte Mège Mouriés, who patented his product in France and Britain in 1869. His process required the softer component from fractionated tallow, skimmed milk, and macerated cow’s udder. The product was described as mixed glycerol esters of oleic and margaric acids and was therefore called oleo-margarine. Margaric acid was thought to be heptadecanoic acid (17:0), but it was actually a eutectic mixture of palmitic (16:0) and stearic (18:0) acids. Even this early process involved fractionation and enzymes. (Both margaric
and margarine should be pronounced with a hard g as in Margaret. All three words come from the Greek word for "pearl"—margarites.

For a long time margarine was considered as a cheap and inferior substitute for butter. In several countries regulations were passed that prohibited the addition of coloring matter so that white margarine would compare even less favorably with the more familiar yellow butter. Now the situation is different. These impediments have disappeared and margarine is widely accepted as having several advantages over butter. It is a more flexible product that can be varied for different markets and modified to meet new nutritional demands, such as desirable levels of cholesterol, phytosterols, saturated or trans acids, and fat content, as well as the statutory levels of certain vitamins. General information on margarine is available in Refs. 10–12. Table margarine is made from appropriate oils and fats (soybean, rapeseed/canola, cottonseed, palm, palm kernel, coconut), which may have been fractionated, blended, hydrogenated in varying degrees, and/or interesterified. Fish oil (hydrogenated or not) may also be included. Other ingredients include surface-active agents, proteins, salt, and water along with preservatives, flavors, and vitamins.

Margarine production involves three basic steps: emulsification of the oil and aqueous phases, crystallization of the fat phase, and plasticification of the crystallized emulsion. Water-in-oil emulsions are cooled in scraped-wall heat exchangers during which time fat crystallization is initiated, a process known as nucleation, and during which the emulsion drop size is reduced. There follows a maturing stage in working units during which crystallization approaches equilibrium, though crystallization may continue even after the product has been packed. The lipid in a margarine is part solid (fat) and part liquid (oil), and the proportion of these two varies with temperature. The solid/liquid ratio at different temperatures is of paramount importance in relation to the physical nature of the product.

Individual crystals are between 0.1 and several micrometers in size, and form clusters or aggregates of 10–30 \( \mu \)m. One gram of fat phase may contain up to \( 10^{12} \) individual crystals. The aqueous phase is present in droplets, generally 2–4 \( \mu \)m in diameter, stabilized by a coating of fat crystals.

It is desirable that margarine taken from the refrigerator at 4°C should spread easily. For this to happen the proportion of solids should be 30–40% at that temperature and should not exceed the higher value. For the sample to "stand up" at room temperature (and not collapse to an oily liquid) it should still have 10–20% solids at 10°C. Finally, so that it melts completely in the mouth and not have a waxy mouthfeel, the solid content should be less than 3% at 35°C. These are important parameters that can be attained with many different fat blends. Formulations have to be changed slightly to make the product suitable for use in hot climates.

Fats usually crystallize in two different forms, known as \( \beta' \) and \( \beta \). Of these the \( \beta \) form is thermodynamically more stable and will therefore be formed in many fats and fat blends. But sometimes the fat remains in the \( \beta' \) form. For margarines and other spreads the \( \beta' \) form is preferred because the crystals are smaller, are able to entrap more liquid to give firm products with good texture and mouthfeel, and impart a high gloss to the product. The \( \beta \) crystals, on the other hand, start small but tend to agglomerate and can trap less liquid. It is therefore desirable to choose a blend of oils that crystallize in the \( \beta' \) form.

Margarines and shortenings made from rapeseed/canola, sunflower, and soybean oil after partial hydrogenation tend to develop \( \beta \) crystals. This can be inhibited
or prevented by the incorporation of some cottonseed oil, hydrogenated palm oil or palm olein, tallow, modified lard, or hydrogenated fish oil, all of which stabilize crystals in the $\beta'$ form. The canola, sunflower, and soybean oils share very high levels of $C_{18}$ acids, whereas the remainder have appreciable levels of $C_{16}$ acids (or other chain length) along with the $C_{18}$ acids and thus contain more triacylglycerols with acids of mixed chain length.

There are many formulations for making margarines, and different recipes are used around the world depending on the oils most readily available in any particular locality. Practically all of them contained partially hydrogenated oils and therefore appreciable levels of trans acids. Attempts are now being made to reduce the levels of such acids on nutritional grounds. Considerable progress has been made in European formulations, and the average content of trans acids has fallen in recent years. Many preparations approach zero trans. New legislation in the United States will make it necessary to indicate the content of trans acids, and this will probably have an influence on formulations. However, it must be realized that it is not possible to make spreads without a proportion of solid triacylglycerols, which must contain saturated and/or trans monoene acids. If the level of trans acids is to fall, then there must be some rise in the content of saturated acids. Nor is it sufficient merely to blend hardstock (material with a high proportion of solid triacylglycerols) with oils containing “healthy” mono- and polyunsaturated acids. The blended oils may have to be interesterified to get the appropriate distribution of fatty acids in the triacylglycerols. It is impossible to list all of the formulations used to make margarines, and the following list is merely indicative (in the following blends, “hydrogenated” means “partially hydrogenated”):

- Blends of hydrogenated soya bean oils with or without unhydrogenated soybean oil
- Blends of canola oil, hydrogenated canola oil, and hydrogenated palm oil or palm stearin
- Blends of various hydrogenated cottonseed oils
- Blends of edible tallow with vegetable oils (soybean, coconut)
- Blends of palm oil with hydrogenated palm oil and a liquid oil (rapeseed, sunflower, soybean, cottonseed, olive)

For hot climates a harder formulation is required, as in the following examples from Malaysia:

- Palm oil (60%), palm kernel oil (30%), and palm stearin (10%)
- Palm stearin (45%), palm kernel oil (40%), and liquid oil (15%)

Table 8 gives details of the fatty acid composition of spreading fats and Table 9 provides information on production levels of margarine in the period 1994–1999.

Margarine is expected to have a shelf life of about 12 weeks. With good ingredients and the absence of prooxidants (e.g., copper), oxidative deterioration is not likely to be a problem. However, care must be taken to avoid microbiological contamination in the aqueous phase. This is avoided by hygienic practices during manufacture, the addition of some salt (8–10% in the aqueous phase, corresponding to slightly more than 1% in the margarine), control of pH of any cultured milk that may be used, and careful attention to droplet size in the emulsion.
Table 8  Approximate Fatty Acid Composition of Spreading Fats (%)

<table>
<thead>
<tr>
<th>Fat</th>
<th>Saturated</th>
<th>Monoene&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Polyene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Butter</td>
<td>63–70</td>
<td>28–31</td>
<td>1–3</td>
</tr>
<tr>
<td>Stick (packet), veg. oils only</td>
<td>18–21</td>
<td>45–66</td>
<td>14–35</td>
</tr>
<tr>
<td>Stick (packet), veg. and animal fats</td>
<td>29–40</td>
<td>46–52</td>
<td>9–19</td>
</tr>
<tr>
<td>Soft (tub)</td>
<td>17–19</td>
<td>35–52</td>
<td>29–48</td>
</tr>
</tbody>
</table>

<sup>a</sup>U.S. stick margarines contain 17–36% trans acids and soft margarines contain 10–18% trans acids. However, products with lower levels of trans acids are now being produced.

The levels of total trans acids (mainly 18:1 but also 18:2 and 18:3) in margarines from various countries are listed in Table 10 (19–23). Levels have declined over the last 10 years and are noticeably lower in Europe than in North America. However, margarines are not the only source of dietary trans fatty acids. Such acids are also obtained from dairy produce and from baked goods made with partially hydrogenated vegetable fats. Ratnayake et al. (22) report that with a trans fatty acid consumption of about 8.4 g/day in Canada, only about 0.96 g (11%) comes from the consumption of margarine. Wolff et al. (24) have drawn attention to the very different profile of trans monoene fatty acids consumed in France and Germany compared with consumption in North America. These differences reflect the differing nature of trans acids from dairy produce on the one hand and industrially hydrogenated vegetable oils on the other.

Margarines are now available with added phytosterols, which, it is claimed, are capable of reducing blood cholesterol levels. The phytosterols are obtained from tall oil and added as hydrogenated sterol esters or from soybean oil and added as unsaturated sterol esters to margarines at around the 8% level. Margarine is a suitable vehicle for phytosterol addition because it is a food used widely and regularly, and is unlikely to be overconsumed. Intake of phytosterols is normally 200–400 mg/day, though higher for vegetarians, and the intake of 1.6–3.3 g/day, recommended by those offering this special margarine, is markedly higher. Normally about 50% of

Table 9  Production of Margarine (MMT Expressed in Terms of Normal Fat Levels and Including Vanaspati), 1994–1999

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>World total</td>
<td>9.00</td>
<td>9.19</td>
<td>9.31</td>
<td>9.51</td>
<td>9.36</td>
<td>9.50</td>
</tr>
<tr>
<td>EU-15</td>
<td>2.31</td>
<td>2.43</td>
<td>2.40</td>
<td>2.52</td>
<td>2.41</td>
<td>2.30</td>
</tr>
<tr>
<td>E. Europe</td>
<td>0.52</td>
<td>0.59</td>
<td>0.59</td>
<td>0.63</td>
<td>0.62</td>
<td>0.63</td>
</tr>
<tr>
<td>Ex-USSR</td>
<td>0.56</td>
<td>0.46</td>
<td>0.47</td>
<td>0.49</td>
<td>0.48</td>
<td>0.61</td>
</tr>
<tr>
<td>USA</td>
<td>1.19</td>
<td>1.13</td>
<td>1.12</td>
<td>1.07</td>
<td>1.05</td>
<td>1.02</td>
</tr>
<tr>
<td>Brazil</td>
<td>0.40</td>
<td>0.42</td>
<td>0.44</td>
<td>0.45</td>
<td>0.47</td>
<td>0.47</td>
</tr>
<tr>
<td>India</td>
<td>0.91</td>
<td>0.96</td>
<td>1.02</td>
<td>1.03</td>
<td>0.96</td>
<td>1.01</td>
</tr>
<tr>
<td>Pakistan</td>
<td>1.33</td>
<td>1.36</td>
<td>1.38</td>
<td>1.41</td>
<td>1.44</td>
<td>1.47</td>
</tr>
<tr>
<td>Turkey</td>
<td>0.56</td>
<td>0.58</td>
<td>0.61</td>
<td>0.63</td>
<td>0.65</td>
<td>0.67</td>
</tr>
<tr>
<td>Other</td>
<td>1.22</td>
<td>1.26</td>
<td>1.28</td>
<td>1.28</td>
<td>1.28</td>
<td>1.32</td>
</tr>
</tbody>
</table>
Table 10  Presence of trans Fatty Acids in Margarines

<table>
<thead>
<tr>
<th>Country (year of publication)</th>
<th>Range (%)</th>
<th>Mean (%)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Germany (1997)</td>
<td>0.2–5</td>
<td>1.5</td>
<td>19</td>
</tr>
<tr>
<td>Belgium, Hungary, and Britain (1996)</td>
<td>1–24</td>
<td>9.7</td>
<td>20</td>
</tr>
<tr>
<td>Denmark (1998), soft</td>
<td>0.4</td>
<td></td>
<td>21</td>
</tr>
<tr>
<td>Denmark (1998), hard</td>
<td>4.1</td>
<td></td>
<td>21</td>
</tr>
<tr>
<td>Canada (1998), tub</td>
<td>1–46*</td>
<td>18.8</td>
<td>22</td>
</tr>
<tr>
<td>Canada (1998), hard</td>
<td>16–44</td>
<td>34</td>
<td>22</td>
</tr>
<tr>
<td>Hawaii (2001), cup</td>
<td>1–19</td>
<td>12.1</td>
<td>23</td>
</tr>
<tr>
<td>Hawaii (2001), carton</td>
<td>18–27</td>
<td>23.4</td>
<td>23</td>
</tr>
</tbody>
</table>

*aMainly 15–20%.

ingested cholesterol is absorbed but with an adequate supply of phytosterols, which are absorbed only at the 5% level, absorption of cholesterol falls to about 20%.

Spreads with reduced levels of fat (40% or less) are popular with consumers (as an alternative to discipline in the amount of normal spread consumed). These spreads contain more water than the full-fat spreads and require emulsifiers (monoacylglycerols or polyglycerol esters). It is also usual to add thickeners, such as gelatin, sodium alginate, pectin, and carrageenan, to the aqueous phase. Industrial margarines are used mainly for bakery products and are discussed in Sec. IV.

D. Vanaspati

It has been estimated that production of vanaspati in 1998 (49) was 4.7 MMT (mainly in Pakistan 1.4, India 1.0, Iran 0.5, Egypt 0.4, and Turkey 0.4). Vanaspati can be considered as vegetable ghee. It is used mainly for frying and for the preparation of sauces, sweets, and desserts. Traditionally, vanaspati was a blend of hydrogenated seed oils (cottonseed, groundnut, soybean, rapeseed/canola and palm), but increasingly palm oil has become a significant component. The product should melt between 31°C and 41°C, though generally it is close to 37°C in India and 36 ± 2°C in Pakistan. A wide range of oils are used, including soybean, rapeseed, sunflower, cottonseed, palm olein, and palm oil. Because of the method of production involving hydrogenation, vanaspati contains high levels of acids with trans unsaturation (more than 50% in India and about 27% in Pakistan). With increasing use of palm oil in vanaspati the need for hydrogenation is reduced with a consequent fall in the level of trans acids. Figures around 3% have been reported in Pakistan (10,25).

IV. BAKING FATS, DOUGHS, SHORTENINGS

The use of oils and fats in baking processes ranks with frying and spreads as a major food use of these materials. The products range from breads and layered doughs to cakes, biscuits (cookies) and biscuit fillings, pie crusts, short pastry, and puff pastry. The fats used to produce this wide range of baked goods vary in their properties and particularly in their melting behavior and plasticity. It is possible to attain these properties with different blends of oils, and preferred mixtures vary in different
regions of the world. In addition to the desired physical properties, it is necessary to meet two further requirements. One is oxidative stability related to the shelf life of the baked goods. The other is the necessity of responding to current nutritional demands. A good baked item will be tasty, will have good texture, will have a reasonable shelf life in terms of rancidity and palatability and texture, and it will be a healthy food. Sometimes the pressure for appropriate physical properties and nutritional requirements work in opposite directions and a compromise has to be made. As already discussed with the spreads, a plastic fat containing solid and liquid components must have some solid triacylglycerols, which implies a certain level of saturated acids or of acids with trans unsaturation despite the nutritional concerns associated with these compounds.

Fats used to make doughs of various kinds are almost entirely plastic fats, i.e., mixtures of solid and liquid components that appear solid at certain temperatures and that deform when a pressure is applied. Fats exert their influence by interaction with the flour and (sometimes) sugar, which are the other major components of a baked product.

Going back to the important physical properties, the solid fat ratio at various temperatures is now usually measured by pulsed $^1$H nuclear magnetic resonance. Plasticity depends on the solid components being in the correct polymorphic form (see Sec. III.C). Tests have also been devised to determine the extent of oxidation and to assess shelf life.

Baking fats may include butter or margarine which are more than 80% fat and also contain an aqueous phase, or they may be shortenings with 100% fat. These are described as shortenings as they give pastry the crispness and flakiness that is suitable for its edible purpose. Industrial margarine has the fat/water ratio required of margarine but differs from margarine spread in that it has fat components that produce the physical properties required by its final end use. Changes in the composition of fat in margarine spread designed to increase their nutritional value have not always carried through to the baking fats, which are often richer in saturated fatty acids and/or acids with trans unsaturation (see Sec. III.C). But there seems little doubt that the appropriate changes will come. Baked goods contain what is described as “hidden” fat, and it is easy to forget the fats they contain when delicious pastries, cakes, and biscuits are being eaten.

The prime function of fat in a cake is to assist in aeration and to modify the texture of the product. The first stage in making a cake is to produce a batter containing a fine dispersion of air bubbles largely stabilized by fat crystals. During baking the fat melts and the water-in-oil emulsion inverts with the air being trapped in the aqueous phase. As baking continues the starch is hydrated and gelatinized, the protein starts to coagulate, and the air cells expand through the presence of steam and carbon dioxide (produced from baking powder).

In short pastry, aeration is only of secondary importance. The fat needs to be fairly firm and should be distributed throughout the dough as a thin film; lard, beef tallow olein, and hardened vegetable oils may be employed. Sometimes butter or margarine is used.

In puff pastry (pie crust, Danish pastries, croissants), fat acts as a barrier separating the layers of dough from one another. Liberation of gas or steam during baking produces a layer structure. This requires a fat of higher melting point fat than normal (about 42°C) with a higher solid fat content achieved through an appropriate
degree of hydrogenation. Small amounts of fat (2–5%) are added to bread dough. Additional information is available in Refs. 26–28.

V. SALAD OILS, MAYONNAISE AND SALAD CREAM, FRENCH DRESSINGS

Salad oils, used to make mayonnaise and salad cream, should be oxidatively stable and free of solids even when stored in a refrigerator at about 4°C. Several vegetable oils may be used. Those containing linolenic acid (soybean oil, rapeseed/canola oil) are usually lightly hydrogenated to enhance oxidative stability. All oils are generally winterized to remove high-melting glycerides that would crystallize, as well also waxes present in solvent-extracted oil. The latter lead to a haze in the oil when it is cooled. Salad oils must pass a “cold test,” which requires that the oil remain clear for 5.5 hours at refrigeration temperature. After appropriate treatment, soybean, rapeseed/canola, corn, and sunflower oils are used to produce mayonnaise.

Mayonnaise is an oil-in-water emulsion containing between 65% (legal minimum) and 80% of oil. The aqueous phase contains vinegar, citric acid, and egg yolk. This last contains lecithin, which serves as an emulsifying agent. Lemon and/or lime juice, salt, syrups, seasonings, spices, and antioxidants are optional constituents. These components may be mixed together at temperatures not exceeding 5°C (cold process) or at temperatures around 70°C (hot process). A typical mayonnaise contains vegetable oil (75–80% by weight), vinegar (9.4–10.8%), egg yolk (7.0–9.0%), and small amounts of sugar, salt, mustard, and pepper (27). “Light” mayonnaise contains about 30–40% of oil, and in low-calorie dressings the level is 10% to less than 3%.

Salad creams are similar but contain much less oil (30–40%) along with cooked starch materials, emulsifiers, and gums to provide stability and thickness. They are cheaper than mayonnaise.

French dressings are temporary emulsions of oil, vinegar, or lemon juice, and seasonings. Because the emulsions are not stable the dressings should be shaken before use. A nonseparating product can be made by addition of egg yolk or other emulsifying agent.

According to Ref. 25, production levels in the United States during the period 1993–1998 were 1.35–1.38 million pounds, comprising salad dressings and mayonnaise (0.90–0.97) and pourable dressings (0.39–0.45).

Additional information is available in Refs. 8, 30, and 31.

VI. CHOCOLATE AND CONFECTIONERY FATS

Chocolate is an important fat-containing food based mainly, but not always entirely, on cocoa butter. Confectionery fats are materials with similar physical/functional properties to cocoa butter. Legal definitions of chocolate limit the amount of fat other than cocoa butter that may be used. The incorporation of milk fat into chocolate, the limited use of other fats, and the complete replacement of cocoa butter are discussed later in this section. The most recent book on this topic was published by Beckett (32).

Annual production of cocoa beans (2.9 MMT), cocoa butter (1.5 MMT), and chocolate (5 MMT) is reported to be at about the levels indicated in parentheses. A second source gives a figure of about 1.7 MMT for cocoa butter. Cocoa beans contain
50–55% fat. Production figures for cocoa butter are not included in the statistics generally cited for oil and fat production (Sec. I.C).

Harvested pods are broken open and left in heaps on the ground for about a week during which time the sugars ferment. The beans are then sun-dried and are ready for transportation and storage. To recover the important components the beans are roasted (~150°C), shells are separated from the cocoa nib, and the latter is ground to produce cocoa mass. When this is pressed it yields cocoa butter and cocoa powder, which still contains some fat. Typically, 100 g of beans produces 40 g of cocoa butter by pressing, expelling, or solvent extraction, 40 g of cocoa powder, and 20 g of waste material (shell, moisture, dirt, etc.). Cocoa powder is the residue after extraction and still contains 10–24% fat. Increasingly the beans are processed in the country where they grow and cocoa liquor, cocoa powder, and cocoa butter (usually in 25-kg parcels) are exported to the chocolate-producing countries (33).

Cocoa butter is a solid fat (mp 32–35°C) obtained from the cocoa bean (*Theobroma cacao*) along with cocoa powder. Both the butter and the powder are important ingredients in chocolate. Cocoa butter is in high demand because its characteristic melting behavior gives it properties that are significant in chocolate. At ambient temperatures it is hard and brittle, giving chocolate its characteristic snap, but also it has a steep melting curve that allows for complete melting at mouth temperature. This gives a cooling sensation and a smooth creamy texture. Typical figures for Ghanaian cocoa butter taken from Table 11 show that the content of solid falls from 45% to 1% between 30°C and 35°C. The hardness of cocoa butter is related to its solid fat content at 20°C and 25°C. This melting behavior is related to the chemical composition of cocoa butter. The fat is rich in palmitic (24–30%), stearic (30–36%), and oleic acids (32–39%), and its major triacylglycerols are of the kind SOS where S represents saturated acyl chains in the 1- and 3-positions and O represents an oleyl chain in the 2-position. There are three major components: POP, POSt, and StOSt (P = palmitic acid and St = stearic acid). Cocoa butter has a high content of saturated acids, which raises health concerns; however, it has been argued that much of this is stearic acid, which is not considered to be cholesterolemic. Cocoa butter is also a rich source of flavonoids, which are considered to be powerful antioxidants (34).

Cocoa is grown mainly in West Africa, Southeast Asia, and South and Central America. The composition of cocoa butter from these different sources varies slightly, as shown in Table 11 for cocoa butter from Ghana, Ivory Coast, Brazil, and Malaysia. There are small differences in fatty acid composition that are reflected in the iodine value and the melting point, but more significantly in the triacylglycerol composition and the melting profile. The content of the important SOS triacylglycerols varies between 87.5% in Malaysian and 71.9% in Brazilian cocoa butter, with the African samples midway between these extremes. There is some evidence that the cocoa butters of different geographic origin are becoming more alike (33).

The crystal structure of cocoa butter has been studied intensively because of its importance in our understanding of the nature of chocolate. The solid fat has been identified in six crystalline forms designated I–VI. Some crystals show double chain length (D) and some triple chain length (L). The six forms have the following melting points (°C) and D/T structure: I (17.3, D), II (23.3, D), III (25.5, D), IV (27.3, D), V (33.8, T), and VI (36.3, T). Of these, form V is the one preferred for chocolate. This crystalline form gives good molding characteristics and has a stable gloss and favorable snap at room temperature. Procedures to promote this form are
Table 11  Composition and Properties of Cocoa Butter from Different Countries (35)

<table>
<thead>
<tr>
<th>Factor</th>
<th>Ghana</th>
<th>Ivory Coast</th>
<th>Brazil</th>
<th>Malaysia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iodine value</td>
<td>35.8</td>
<td>36.3</td>
<td>40.7</td>
<td>34.2</td>
</tr>
<tr>
<td>Melting point (°C)</td>
<td>32.2</td>
<td>32.0</td>
<td>32.0</td>
<td>34.3</td>
</tr>
<tr>
<td>Diacylglycerols</td>
<td>1.9</td>
<td>2.1</td>
<td>2.0</td>
<td>1.8</td>
</tr>
<tr>
<td>Free acid (%)</td>
<td>1.53</td>
<td>2.28</td>
<td>1.24</td>
<td>1.21</td>
</tr>
<tr>
<td>Component acids</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Palmitic</td>
<td>24.8</td>
<td>25.4</td>
<td>23.7</td>
<td>24.8</td>
</tr>
<tr>
<td>Stearic</td>
<td>37.1</td>
<td>35.0</td>
<td>32.9</td>
<td>37.1</td>
</tr>
<tr>
<td>Oleic</td>
<td>33.1</td>
<td>34.1</td>
<td>37.4</td>
<td>33.2</td>
</tr>
<tr>
<td>Linoleic</td>
<td>2.6</td>
<td>3.3</td>
<td>4.0</td>
<td>2.6</td>
</tr>
<tr>
<td>Arachidic</td>
<td>1.1</td>
<td>1.0</td>
<td>1.0</td>
<td>1.1</td>
</tr>
<tr>
<td>Component triacylglycerols</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trisaturated</td>
<td>0.7</td>
<td>0.6</td>
<td>trace</td>
<td>1.3</td>
</tr>
<tr>
<td>Monounsaturated</td>
<td>84.0</td>
<td>82.6</td>
<td>71.9</td>
<td>87.5</td>
</tr>
<tr>
<td>POP</td>
<td>15.3</td>
<td>15.2</td>
<td>13.6</td>
<td>15.1</td>
</tr>
<tr>
<td>POSl</td>
<td>40.1</td>
<td>39.0</td>
<td>33.7</td>
<td>40.4</td>
</tr>
<tr>
<td>StOsSt</td>
<td>27.5</td>
<td>27.1</td>
<td>23.8</td>
<td>31.0</td>
</tr>
<tr>
<td>Diunsaturated</td>
<td>14.0</td>
<td>15.5</td>
<td>24.1</td>
<td>10.9</td>
</tr>
<tr>
<td>Polynsaturated</td>
<td>1.3</td>
<td>1.3</td>
<td>4.0</td>
<td>0.3</td>
</tr>
<tr>
<td>Solid content (pulsed NMR)—</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>tempering 40 hours at 26°C</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20°C (%)</td>
<td>76.0</td>
<td>75.1</td>
<td>62.6</td>
<td>82.6</td>
</tr>
<tr>
<td>25°C (%)</td>
<td>69.6</td>
<td>66.7</td>
<td>53.3</td>
<td>77.1</td>
</tr>
<tr>
<td>30°C (%)</td>
<td>45.0</td>
<td>42.8</td>
<td>23.3</td>
<td>57.7</td>
</tr>
<tr>
<td>35°C (%)</td>
<td>1.1</td>
<td>0.0</td>
<td>1.0</td>
<td>2.6</td>
</tr>
</tbody>
</table>

Adapted from Ref. 35. The original paper contains more details along with information on cocoa butter from India, Nigeria, and Sri Lanka.

necessary, and its change to form VI must be inhibited. Form V is usually obtained as a result of extensive tempering, i.e., putting molten chocolate through a series of cooling and heating processes that have been found to optimize production of the appropriate polymorph. Alternatively, molten chocolate can be seeded with cocoa butter already in form V.

Transition from form V to the more stable form VI leads to the appearance of white crystals of fat on the surface of the chocolate. This phenomenon is termed “bloom.” It is promoted by fluctuations in temperature during storage and by migration of liquid oils from nut centers. It is a harmless change but is considered undesirable because it may be mistaken for microbiological contamination. Bloom can be inhibited by addition of a little 2-oleo 1,3-dibehenin (BOB) to the cocoa butter. This phenomenon is discussed in more detail by Smith (36), Padley (37), and Hammond (38).

The simplest plain chocolate contains sugar and cocoa liquor, with cocoa butter the only fat present. Padley (37) reports that a typical plain chocolate contains cocoa mass (~40%, which contains some cocoa butter), sugar (~48%), added cocoa butter (~12%), and small amounts of lecithin and other materials. The total fat content of this mix is about 31%. In some European countries is is permissible to replace cocoa
butter with up to 5% of another fat with similar fatty acid and triacylglycerol composition taken from a prescribed list of tropical fats (39,40). This represents about 15% of the fat phase. The permitted tropical fats come from palm, illipe, shea, sal, kokum, and mango and may be used in a fractionated form (Table 12).

Milk chocolate contains between 3.5% and 9% of milk fat, and white chocolate is based on sugar and cocoa liquor and cocoa butter. If the latter is not entirely refined it will retain some of the flavor normally associated with chocolate. Chocolate normally contains up to 0.4% of lecithin, usually from soybeans. This aids the processing of the chocolate by reducing the viscosity of molten chocolate. Polyglycerol ricinoleate is sometimes added to to optimize viscosity.

Cocoa butter alternatives (CBAs) is a general name covering cocoa butter equivalents (CBEs), cocoa butter improvers (CBIs), cocoa butter replacers (CBRs) and cocoa butter substitutes (CBSs) (36–38).

CBEs have the same general chemical composition and hence the same physical properties as cocoa butter and include the tropical oils described above and sometimes designated as hard butters. These can be blended to give mixtures of POP, POSl, and StOSl very similar to cocoa butter and fully miscible with it. The level at which cocoa butter can be replaced by a CBE is limited on a legal basis and not on a functional basis. CBEs must be compatible with cocoa butter by virtue of their similar fatty acid and triacylglycerol composition, have a melting range equivalent to that of cocoa butter, yield the β polymorph when processed and tempered in the same way as cocoa butter, and give a product that is at least as good as cocoa butter with respect to bloom. The market for CBEs in those European countries where their use in chocolate is permitted is estimated to be 20,000–25,000 metric tonnes, but it could rise to three times this level if all European Union (EU) countries accepted their use as legal.

CBRs are usually based on vegetable oils (soybean, cottonseed, palm) that have been fractionated and partially hydrogenated. They contain trans unsaturated acids at levels up to 60% and have a different triacylglycerol composition from cocoa butter. They do not require tempering but should be compatible with cocoa butter.

CBSs are usually based on lauric fats. They share some of the physical properties of cocoa butter but have a different composition. Coatings based on CBS fats do not require to be tempered but are used in the molten state for enrobing. They give a superior gloss and have very sharp melting characteristics. Further information is given in Refs. 41–43.

Table 12  Typical Triacylglycerol Composition of Cocoa Butter and Some Fractions Used as Permitted Partial Replacers

<table>
<thead>
<tr>
<th></th>
<th>Cocoa butter</th>
<th>Palm midfraction</th>
<th>Illipe</th>
<th>Shea stearin</th>
<th>Sal stearin</th>
<th>Mango kernel stearin</th>
</tr>
</thead>
<tbody>
<tr>
<td>POP</td>
<td>16</td>
<td>43</td>
<td>7</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>POSl</td>
<td>38</td>
<td>8</td>
<td>31</td>
<td>10</td>
<td>13</td>
<td>15</td>
</tr>
<tr>
<td>StOSl</td>
<td>23</td>
<td>1</td>
<td>50</td>
<td>66</td>
<td>64</td>
<td>65</td>
</tr>
<tr>
<td>Total</td>
<td>77</td>
<td>52</td>
<td>88</td>
<td>77</td>
<td>60</td>
<td>83</td>
</tr>
</tbody>
</table>

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VII. **ICE CREAM**

The annual production of ice cream in the United States is reported to be about 54 million hectoliters (i.e., 5400 million liters), suggesting that the global figure is at least twice this level. This quantity of ice cream will contain around 0.8–1.0 MMT of fat, which will be mainly from milk fat but include a range of vegetable fats such as sunflower, groundnut, palm, palm kernel, and coconut.

Ice cream contains water (60–70%) and total solids (30–40%), with the latter including fat (5–12%), milk solids other than fat (10–12%), sucrose (12–14%), glucose solids (2–4%), emulsifier (0.2–0.5%), and stabilizer (0.1–0.3%). Legal requirements for fat vary from country to country as does the possibility of replacing some or all of the dairy fat with vegetable fat.

Fat in ice cream contributes to structure. It stabilizes the aerated foam, improves melting resistance, imparts creaminess, and contributes to taste. Its most important properties are melting characteristics, solid-to-liquid ratio at various temperatures, and its taste profile.

Production of ice cream occurs through nine stages: selection and weighing of ingredients, mixing of these in an appropriate sequence at 20–35°C, pasteurization (70–75°C) or sterilization (95°C), homogenization at 75°C, cooling to less than 5°C, ageing at 5°C for at least 4 hours, freezing to −5 to −10°C, hardening at −25 to −35°C, and storage at −18 to −20°C (44,45).

VIII. **INCORPORATION OF VEGETABLE OILS INTO DAIRY PRODUCTS**

Vegetable oils may be incorporated into dairy products as a replacement for dairy fat. This happens when local supplies are inadequate as in some tropical countries where the climate is not suitable for large-scale dairy farming and also for consumers concerned about the saturated acids and cholesterol present in milk fat. In addition, it is possible to produce milk fat replacements in a more convenient form as, for example, in long-life cream. The possible use of vegetable fat in ice cream has already been discussed in Sec. VII.

So-called filled milk is made from skim milk powder reconstituted with an appropriate vegetable oil. This should be free of linolenic acid, have a low content of linoleic acid, and contain antioxidant so that it is oxidatively stable. Palm oil, palm kernel oil, and coconut oil are most frequently used, and these may be partially hydrogenated to provide further stability against oxidation.

Nondairy coffee whiteners, available in powder or liquid form, generally contain 35–45% fat, which is usually partially hydrogenated palm kernel oil.

Cheeses have been developed based on vegetable fat rather than dairy fat. A range of formulations have been described incorporating soybean oil, with or without hydrogenation, palm oil, rapeseed oil, lauric oils, and high-oleic sunflower oil. Attempts have been made to incorporate into these products some of the short chain acids that are characteristic of milk fat and give cheese some of its characteristic flavor (46,47).

Nondairy whipping creams made with hardened palm kernel oil and coconut oil (each about 17%) are convenient because they have a long shelf life at ambient temperature. They are popular in Britain. First produced for the bakery and catering
market with high overrun and good shape retention, they are now supplied to the retail market for domestic use. Pouring creams containing about 9% of each of the two lauric oils are also available. Both creams also contain buttermilk powder (7%), guar gum (0.10–0.15%), emulsifying agent (0.30–0.35), β-carotene (0.25%), and water.

IX. EDIBLE COATINGS AND SPRAY PROCESSING

Foods are sometimes coated with thin layers of edible material to extend shelf life by minimizing moisture loss, to provide gloss for aesthetic reasons, and to reduce the complexity and cost of packaging. The thin layers may be carbohydrate, protein, lipid, or some combination of these. The lipids most commonly used are waxes (candelilla, carnauba, or rice bran), appropriate triacylglycerols, or acetylated monoacylglycerols. The latter are capable of producing flexible films at temperatures below those appropriate for the waxes even though they are poorer moisture barriers. The foods most frequently coated are citrus fruits (oranges and lemons), deciduous fruits (apples), vegetables (cucumbers, tomatoes, potatoes, candies and confectioneries, nuts, raisins, cheeses, and starch-based products (cereals, doughnuts, and ice cream cones and wafers).

Vegetable oils used to coat food products must be liquid at room temperature and must have high oxidative stability. They serve as a moisture barrier, a flavor carrier, a lubricant or release agent, as an antidust or anticake agent, and as a gloss enhancer. They are used at low levels and are sprayed onto large exposed surfaces of products during roasting frying, or handling. Traditionally they are made from commodity oils like soybean or cottonseed. These are cheap but require elaborate processing (partial hydrogenation and fractionation) to develop the required physical state and chemical stability. New high-oleic oils may also be used. These are more costly but they will bring added value in terms of their superior nutritional properties resulting from lower trans acids and lower saturated acids and in the reduced need for processing. Lauric oils, such as coconut oil, palm kernel oil, are used to spray cracker-type biscuits to provide an attractive appearance, maintain crispness by acting as a barrier to moisture, and improve eating quality (50).

X. EMULSIFYING AGENTS

Fatty acids and their derivatives are amphiphilic. This means that their molecules have hydrophilic (lipophobic) and lipophilic (hydrophobic) regions. If these are appropriately balanced, then the molecules can exist in a physically stable form between aqueous and fatty substances. They can therefore be used to stabilize both oil-in-water and water-in-oil emulsions and are important components of many of the fat-based products that have been described in the earlier sections of this chapter. Applications of emulsifiers in foods include film coatings, stabilizing and destabilizing emulsions, modification of fat crystallization, dough strengthening, crumb softening, and texturization of starch-based foods. Krog (51) estimates that production of food emulsifiers is about 250,000 metric tonnes of which about 75% is monoacylglycerols or compounds derived from these.

Monoacylglycerols are most often made by glycerolysis of natural triacylglycerol mixtures in the presence of an alkaline catalyst (180–230°C, 1 hour). Fat and
glycerol (30% by weight) will give a mixture of monoacylglycerols (around 58%, mainly the 1-isomer), diacylglycerols (about 36%), and triacylglycerols (about 6%). This mixture can be used in this form or it can be subjected to high-vacuum thin-film molecular distillation to give a monoacylglycerol product (around 95% and at least 90% of the 1-monoester) with only low levels of diacylglycerols, triacylglycerols, and free acids. Attempts are being made to develop an enzyme-catalyzed glycerolysis reaction that occurs under milder reaction conditions. The oils most commonly used include lard, tallow, soybean, cottonseed, sunflower, palm, and palm kernel oil—all in hydrogenated or nonhydrogenated form. Glycerol monostearate (GMS) is a commonly used product of this type.

The properties desired in a monoacylglycerol for some specific use may be improved by acylation of one of the free hydroxyl groups by reaction with acid (lactic, citric) or acid anhydride (acetic, succinic, diacetyltartric). For the most part these have the structures shown:

\[
\text{CH}_3(\text{CH}_2)_n\text{COOCHCH(OH)CHOCOR, where R is:}
\]

- \(\text{CH}_3(\text{acetate})\)
- \(\text{CH(OH)COOH (lactate)}\)
- \(\text{CH}_2\text{CH}_2\text{COOH (succinate)}\)
- \(\text{CHOAcCHOacCOOH (diacetyltartrate)}\)
- \(\text{CH}_3\text{C(OH,COOH)CH COOH (citrate)}\)

Propylene glycol (\(\text{CH}_3\text{CHOHCH}_2\text{OH}\)) also reacts with fatty acids to give mixtures of mono (about 55%, mainly 1-acyl) and diacyl esters (about 45%). A 90% monoacyl fraction can be obtained by molecular distillation.

Other compounds include the partial esters of polyglycerols (a polyether with 2–10 glycerol units but mainly 2–4 units), sorbitan and its polyethylene oxide derivatives, the 6-monoacylate sucrose, and stearoyl lactate, usually as the sodium or calcium salt (51, 52).

REFERENCES


I. INTRODUCTION

Lipid biotechnology broadly covers the areas outlined in Scheme 1. Genetic engineering of oilseeds and oil-bearing fruits for improved agronomic properties and altered fatty acid and lipid composition is a rapidly expanding area of lipid biotechnology (1). Plant genetic engineering of edible oilseed crops is covered in Chapter 30 of this book.

This chapter will initially cover the use of microorganisms, such as microalgae, yeasts, molds, and bacteria, for the production of oils and fats containing triacylglycerols and other lipids of commercial interest from nonlipidic and lipid-containing carbon sources. This will be followed by biotransformation of fats, oils, and their constituent fatty acids using whole microbial cells or enzymes isolated from various organisms for the production of fat-based added-value products, as outlined in Scheme 2. A bulk of this section will be devoted to application of triacylglycerol lipases and phospholipases for the preparation of specialty products. Finally several examples will be given in this chapter on the applications of enzymes in the processing of oilseeds and other oil-bearing materials as well as the use of enzymes in the processing of fats and oils.

II. MICROBIAL PRODUCTION OF FATS AND OTHER LIPIDS

A. Fats and Lipids from Biomass of Microorganisms Using Nonlipidic or Lipid-Containing Carbon Sources—Single-Cell Oils

Some microorganisms, such as eucaryotic yeasts, molds, and algae, are known to produce triacylglycerols in their biomass (single-cell oils, SCOs) similar to plant oils,
**Lipid Biotechnology**

<table>
<thead>
<tr>
<th>Oil Plants</th>
<th>Oilseeds/ Oil-bearing fruits</th>
<th>Carbohydrates/ hydrocarbons oils/fats</th>
</tr>
</thead>
</table>

**Genetic engineering**

| Oil extraction/ processing using enzymes (cellulase, phospholipase) | Biotransformation/ biomass production using whole microbial enzymes cells (yeasts, fungi, bacteria, oxygenases, etc.) |

**Transgenic plants - oils with altered fatty acid/ lipid composition**

| Oils/fats | Single cell oils and fat-based added value products for food and non-food applications |

**Scheme 1** Major areas of lipid biotechnology.

**Scheme 2** General principles of modification of fats and other lipids by biotransformation using whole microbial cells or enzymes isolated from various organisms.
Table 1  Lipid Content and Levels of Major Constituent Fatty Acids in Biomass of 
Lipid-Accumulating Microalgae

<table>
<thead>
<tr>
<th>Organism</th>
<th>Lipid content (% w/w of biomass)</th>
<th>Fatty acid (% w/w of total)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>16:0</td>
<td>16:1ω9</td>
</tr>
<tr>
<td>Chlorella vulgaris</td>
<td>39</td>
<td>16</td>
</tr>
<tr>
<td>Botryococcus braunii</td>
<td>53–70</td>
<td>12</td>
</tr>
<tr>
<td>Scenedesmus acutus</td>
<td>26</td>
<td>15</td>
</tr>
<tr>
<td>Navicula pelliculosa</td>
<td>22–32</td>
<td>21</td>
</tr>
</tbody>
</table>

*Fatty acids are designated by number of C atoms (left of colon) and number of cis double bonds (right of colon).

Source: Ref. 2.

whereas prokaryotic organisms, such as bacteria, produce more specific lipids, e.g., wax esters, polyesters, poly-β-hydroxybutyrate, etc. This section will cover a few selected microorganisms that have some potentials of application for the production of triacylglycerols and other lipids from nonlipidic and lipid-containing carbon sources. For a more comprehensive review on this subject, the reader is referred to (2–7).

1. Oils and Fats

Lipid content and the levels of major constituent fatty acids in the biomass of a few selected species of lipid accumulating microorganisms are given in Table 1 for microalgae, Table 2 for yeasts, and Table 3 for molds (fungi).

Table 2  Lipid Content and Levels of Major Constituent Fatty Acids in Biomass of 
Lipid-Accumulating Yeasts

<table>
<thead>
<tr>
<th>Organism</th>
<th>Carbon source</th>
<th>Lipid content (% w/w of biomass)</th>
<th>Fatty acid (% w/w of total)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>16:0</td>
<td>16:1</td>
</tr>
<tr>
<td>Candida curvata D</td>
<td>Whey (lactose)</td>
<td>58</td>
<td>32</td>
</tr>
<tr>
<td>(Apiotrichum curvatum)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cryptococcus albidus</td>
<td>Glucose, ethanol</td>
<td>65</td>
<td>16</td>
</tr>
<tr>
<td>Lipomyces lipofer</td>
<td>Ethanol, glucose</td>
<td>64</td>
<td>37</td>
</tr>
<tr>
<td>Lipomyces starkeyi</td>
<td>Ethanol, glucose</td>
<td>63</td>
<td>34</td>
</tr>
<tr>
<td>Rhodosporidium toruloides</td>
<td>Glucose, mixed sugars</td>
<td>66</td>
<td>18</td>
</tr>
<tr>
<td>Trichosporon pullulans</td>
<td>Ethanol, glucose</td>
<td>65</td>
<td>15</td>
</tr>
<tr>
<td>Rhodotorula glutinis</td>
<td>Glucose, mixed sugars</td>
<td>72</td>
<td>37</td>
</tr>
<tr>
<td>Yarrowia lipolytica</td>
<td>Glucose</td>
<td>32–36</td>
<td>11</td>
</tr>
<tr>
<td>(Candida lipolytica)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Fatty acids are designated by number of C atoms (left of colon) and number of cis double bonds (right of colon).

Source: Ref. 2.
Table 3  Lipid Content and Levels of Major Constituent Fatty Acids in Biomass of Lipid-Accumulating Molds Grown on Glucose as Carbon Source

<table>
<thead>
<tr>
<th>Organism</th>
<th>Lipid content (% w/w of biomass)</th>
<th>Fatty acid (% w/w of total)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>16:0</td>
<td>16:1</td>
</tr>
<tr>
<td>Aspergillus terreus</td>
<td>57</td>
<td>23</td>
</tr>
<tr>
<td>Claviceps purpurea*</td>
<td>31–60</td>
<td>23</td>
</tr>
<tr>
<td>Mucor ramannianus</td>
<td>56</td>
<td>19</td>
</tr>
<tr>
<td>Tolyposporium ehrenbergii</td>
<td>41</td>
<td>7</td>
</tr>
</tbody>
</table>

*Fatty acids are designated by number of C atoms (left of colon) and number of cis double bonds (right of colon).

Some microalgae, as opposed to macroalgae (seaweeds, phytoplankton, etc.), are known to produce substantial amounts of lipids with widely varying fatty acid composition in their biomass (Table 1). Due to relatively high cost of growing algae, these organisms are being considered for the production of specialty lipids to be discussed later, rather than common oils and fats for food uses.

Biomasses of oleaginous yeasts, listed in Table 2, contain high levels of lipids, predominantly triacylglycerols, having fatty acid composition resembling several edible fats and oils (8). Oleaginous yeasts can grow well on a wide variety of substrates, such as pure sugars (e.g., glucose, sucrose, and fructose), mixed sugars contained in molasses, lactose contained in whey, ethanol, and so forth. Whey is a soluble liquor by-product of the cheese-making industry that contains up to 5% w/v lactose in addition to noncoagulated protein. The protein can be separated from lactose by ultrafiltration, and the resulting permeate containing lactose serves as an excellent substrate for the oleaginous yeast, Candida curvata D (= Apiotrichum curvatum ATCC 20509). The fat in the resulting biomass has fatty acid composition and triglyceride structure resembling the natural cocoa butter (5) that fetches a high price. Use of Candida curvata D for the production of cocoa butter substitute from whey permeate is being considered as a commercial process.

Some oleaginous molds synthesize high proportions of lipids, predominantly triacylglycerols, in their biomass using glucose as carbon source (Table 3). Fatty acid composition of the lipids of different molds varies widely (6). Whereas some molds, e.g., Aspergillus terreus and Tolyposporium ehrenbergii, produce single-cell oils having fatty acid composition similar to that of edible plant oils, others produce substantial proportions of less common fatty acids (Table 3). For example, Claviceps purpurea produces large proportions of ricinoleic acid, and some Mucor species produce substantial proportions of γ-linolenic acid (Table 3). Use of such microorganisms for the production of specific polyunsaturated fatty acids will be discussed in Sec. II.A.2.

Bacteria that have been reported to produce substantial proportions of triacylglycerols in their biomass include Actinomycetales, especially mycobacteria, corynebacteria, and Nocardia (9). The biomass of an Arthrobacter sp. contains as much...
as 80% lipids of which about 90% are triacylglycerols (10). However, cell yields of such organisms are rather low, and the presence of potentially toxic lipids in their biomass restricts the usefulness of such microorganisms as producer of triglyceride-rich single-cell oils.

2. Polyunsaturated Fatty Acids

Polyunsaturated fatty acids of the ω3 and ω6 series, which are precursors of eicosanoids, are of great interest in specialty products, such as “Nutraceuticals,” due to their ability to modulate the metabolism of eicosanoids, which play a great role in health and disease (see Chapters 17–20 in Part IV of this book).

Several species of microalgae are known to produce large proportions of ω3 and ω6 polyunsaturated fatty acids and eicosanoids in their biomass (11–13 and Table 4). Thus, a docosahexaenoic acid-containing single-cell oil (DHASCO) containing about 40% docosahexaenoic acid (22:6 ω3) is produced commercially from the biomass of the heterotrophic, nonphotosynthetic marine algae, Cryptothecodinium cohnii (14,15).

Selected fungi that have been reported to produce ω3 and ω6 polyunsaturated fatty acids in their biomass are listed in Table 5. Especially, Mucor javanicus (5), Mortierella isabellina and other Mortierella species (8,30,31) have been used for the commercial production of oils containing γ-linolenic acid (18:3 ω6) as substitute for rather expensive plant oils, such as those from seeds of borage (Borago officinalis), evening primrose (Oenothera biennis), black currant, and other Ribes spp. that also contain 10–25% γ-linolenic acid. Typically, a commercially produced oil from the biomass of a Mortierella sp. contains as major constituent fatty acids palmitic (27%), stearic (6%), oleic (44%), linoleic (12%), and γ-linolenic acid (8%) (30). Optimization of culture conditions of a Mortierella ramanniana species has yielded an oil containing 18% γ-linolenic acid (32). Similarly, commercially produced anachidonic acid-containing single-cell oil (ARASCO) from the biomass of Mortierella alpina contains as much as 40% arachidonic acid (20:4 ω6) (33). Recently, culture conditions have been optimized for the production of an oil containing about 40% dihomogamma-linolenic acid (20:3 ω6) from the biomass of Mortierella alpina (34).

### Table 4  Levels of Major Constituent Polyunsaturated Fatty Acids in Biomass of Lipid-Accumulating Microalgae

<table>
<thead>
<tr>
<th>Organism</th>
<th>Major polyunsaturated fatty acid</th>
<th>(%) w/w of total fatty acids</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Spirulina platensis</em></td>
<td>18:3ω6</td>
<td>21</td>
</tr>
<tr>
<td><em>Dunaliella tertiolecta</em></td>
<td>18:3ω6</td>
<td>32</td>
</tr>
<tr>
<td><em>Porphyridium cruentum</em></td>
<td>20:4ω6</td>
<td>60</td>
</tr>
<tr>
<td><em>Chlorella minutissima</em></td>
<td>20:5ω3</td>
<td>45</td>
</tr>
<tr>
<td><em>Navicula saprophilla</em></td>
<td>20:5ω3</td>
<td>22</td>
</tr>
</tbody>
</table>

*Fatty acids are designated by number of C atoms (left of colon) and number of cis double bonds (right of colon).  
Source  Ref. 2.
Table 5  Levels of Major Constituent Polyunsaturated Fatty Acids in Biomass of Lipid-Accumulating Molds

<table>
<thead>
<tr>
<th>Organism</th>
<th>Fatty acid</th>
<th>(% w/w of total fatty acids)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mortierella isabellina</td>
<td>18:3</td>
<td>8</td>
<td>2</td>
</tr>
<tr>
<td>Mucor javanicus</td>
<td>18:3</td>
<td>15–18</td>
<td>2</td>
</tr>
<tr>
<td>Mucor ambiguus</td>
<td>18:3</td>
<td>11–14</td>
<td>16</td>
</tr>
<tr>
<td>Mortierella rammaniana</td>
<td>18:3</td>
<td>26</td>
<td>17</td>
</tr>
<tr>
<td>Mortierella alpina</td>
<td>20:3</td>
<td>23</td>
<td>18</td>
</tr>
<tr>
<td></td>
<td>20:4</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>Mortierella alpina</td>
<td>20:4</td>
<td>69–79</td>
<td>19</td>
</tr>
<tr>
<td>Mortierella elongata</td>
<td>20:4</td>
<td>15</td>
<td>20</td>
</tr>
<tr>
<td>Mortierella alpina</td>
<td>20:4</td>
<td>31</td>
<td>21</td>
</tr>
<tr>
<td>Mortierella alpina</td>
<td>20:4</td>
<td>43–66</td>
<td>22</td>
</tr>
<tr>
<td>Mortierella alpina</td>
<td>20:4</td>
<td>32–57</td>
<td>23</td>
</tr>
<tr>
<td>Mortierella alpina</td>
<td>20:5</td>
<td>15</td>
<td>24</td>
</tr>
<tr>
<td>Pythium irregulare</td>
<td>20:5</td>
<td>25</td>
<td>25</td>
</tr>
<tr>
<td>Pythium irregulare</td>
<td>20:5</td>
<td>24</td>
<td>26</td>
</tr>
<tr>
<td>Thraustochytrium aureum</td>
<td>20:6</td>
<td>40–50</td>
<td>27</td>
</tr>
<tr>
<td>Thraustochytrium aureum</td>
<td>20:6</td>
<td>29</td>
<td>28</td>
</tr>
<tr>
<td>Thraustochytrium aureum</td>
<td>20:6</td>
<td>40</td>
<td>29</td>
</tr>
</tbody>
</table>

*Fatty acids are designated by number of C atoms (left of colon) and number of cis double bonds (right of colon).

A few species of marine bacteria have also been found to produce substantial proportions of polyunsaturated fatty acids, such as eicosapentaenoic acid (35,36) and docosahexaenoic acid (37) in the lipids of their biomass.

3. Wax Esters

Several reports are known on the production of wax esters by *Acinetobacter* sp. H01-N grown on a wide variety of *n*-alkanes of chain lengths ranging from C₁₆ to C₂₀ (38,39). In general, the chain lengths of the waxes formed depend on the chain length of the alkane (38), e.g.,

\[
C_n \text{ } n\text{-alkane} \rightarrow C_2^n + C_{2n-2} + C_{2n-4} \text{ wax esters}
\]

In addition to saturated wax esters, those containing up to two double bonds are also formed from *n*-alkanes and carbon sources, such as ethanol, acetic acid, propanol, and propionic acid (39). Typical composition of wax esters produced by *Acinetobacter* sp. H01-N from various carbon sources is summarized in Table 6.

The microalgae *Euglena gracilis* (ATCC 12716), grown on yeast malt extract, synthesize wax esters having saturated even carbon numbered acids and alcohols ranging from C₁₂–C₁₈ as the major constituents (40).
Table 6  Wax Esters Formed by *Acinetobacter* sp. H01-N from Various Substrates

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Wax esters(^a)</th>
<th>Fatty acids</th>
<th>Fatty alcohols</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>n</em>-Hexadecane</td>
<td>32:0 + 32:1 + 32:2</td>
<td>16:0 + 16:1</td>
<td>16:0 + 16:1</td>
</tr>
<tr>
<td></td>
<td>36:0 + 38:0 + 40:0</td>
<td>16:0 + 16:1</td>
<td></td>
</tr>
<tr>
<td><em>n</em>-Eicosane</td>
<td>36:1 + 38:1 + 40:1</td>
<td>18:0 + 18:1</td>
<td>20:0 + 20:1</td>
</tr>
<tr>
<td></td>
<td>36:2 + 38:2 + 40:2</td>
<td>20:0 + 20:1</td>
<td></td>
</tr>
<tr>
<td>Ethanol and acetic acid</td>
<td>32:0 + 34:0 + 36:0</td>
<td>16:0 + 16:1</td>
<td>16:0 + 16:1</td>
</tr>
<tr>
<td></td>
<td>32:1 + 34:1 + 36:1</td>
<td>18:0 + 18:1</td>
<td>18:0 + 18:1</td>
</tr>
<tr>
<td></td>
<td>32:2 + 34:2 + 36:2</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^a\)Wax esters, fatty acids, and fatty alcohols are designated by number of C atoms (left of colon) and number of cis double bonds (right of colon).  
Source Ref. 39.

4. Biosurfactants

Yeasts, molds, and bacteria are known to produce a wide variety of extracellular glycolipids and other surface-active substances, which are termed biosurfactants (4,41,42).

Table 7 summarizes the microorganisms and the carbon sources used for the production of several important biosurfactants.

Two common forms of sophorolipids formed from glucose and palm oil are composed of a glycoside of the disaccharide sophorose in which 17-hydroxystearic acid is bound glycosidically at the hydroxyl group or as a lactone in which 17-hydroxystearic acid is bound to sophorose both glycosidically at the hydroxyl group and as an ester at the carboxyl end (Fig. 1). Sophorolipids from some microorganisms contain 13-hydroxydocosanoate as the constituent hydroxy acid (4).

A few common forms of rhamnolipids (Fig. 2) obtained from soybean oil as carbon source consist of rhamnose bound glycosidically to 3-hydroxy fatty acids and their estolides.

Cellobiose lipids produced microbially from coconut oil as carbon source (Fig. 3) contain fatty acids or 3-hydroxy fatty acids esterified at the 2-position of cellobiose and 15,16-dihydroxypalmitic acid bound glycosidically to cellobiose.

Table 7  Microbial Glycolipids as Biosurfactants

<table>
<thead>
<tr>
<th>Glycolipid</th>
<th>Microorganism</th>
<th>Carbon source</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sophorolipids</td>
<td><em>Torulopsis bombicola</em></td>
<td>Glucose/palm oil</td>
<td>43</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Glucose/safflower oil</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Oleic acid</td>
<td>45</td>
</tr>
<tr>
<td></td>
<td><em>Candida apicola</em></td>
<td>Glucose</td>
<td>46</td>
</tr>
<tr>
<td>Rhamnolipids</td>
<td><em>Pseudomonas</em> sp.</td>
<td>Alkanes/glycerol</td>
<td>47</td>
</tr>
<tr>
<td>Trehalose lipids</td>
<td><em>Rhodococcus erythropolis</em></td>
<td>Alkanes</td>
<td>48</td>
</tr>
<tr>
<td></td>
<td><em>Arthrobacter</em> sp.</td>
<td>Alkanes</td>
<td>49</td>
</tr>
<tr>
<td>Cellobiose lipids</td>
<td><em>Ustilago maydis</em></td>
<td>Coconut oil</td>
<td>50</td>
</tr>
</tbody>
</table>

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Figure 1  Common forms of microbial sophorolipids. (From Ref. 4.)

Figure 2  Common forms of microbial rhamnolipids. (From Ref. 4.)

Figure 3  Common forms of microbial cellobiose lipids. (From Ref. 4.)
Some other biosurfactants include emulsan, a lipopolysaccharide produced by an Acinetobacter strain (51), surfactin, produced by *Bacillus subtilis*, which is a heptapeptide linked with β-hydroxyymyristic acid by an ester and an amide bond (52) and penta- and disaccharide lipids produced by *Nocardiad corynebacteroides* sp. (53). Glycolipids containing sophorose and ω- or (ω-1)-hydroxy C_{16}/C_{18} acids, bound as glycosides or esters, have been produced by the yeast *Candida bombicola* grown on glucose and 2-dodecanol (54).

### B. Biotransformation of Fats and Lipids Using Whole Microbial Cells

Bioconversions of fats using whole microbial cells is a promising approach for large-scale economical production of fat-based products. The following selected examples show the applications of microbial biotransformations for the preparation of specific products from fats, fatty acids, and their derivatives.

1. **Wax Esters**

Cells of the alga *Euglena gracilis* (55,56) and other algae (57), as well as microorganisms such as *Corynebacterium* sp. (58), have been shown to catalyze the esterification of fatty acids with long chain alcohols yielding wax esters (Fig. 4).

2. **Hydroxy Acids and Other Oxygenated Fatty Acids**

Numerous examples of microbial oxidation of unsaturated fatty acids for the preparation of hydroxy fatty acids have become known (see Ref. 59 for review). The microbial enzymes involved in oxidation of unsaturated fatty acids to hydroxy acids via hydration (Fig. 5) are termed monooxygenases, which catalyze the incorporation of one atom of molecular oxygen into a substrate, while the other atom is reduced to water (60). Dioxygenases catalyze the incorporation of both atoms of molecular oxygen into the substrate.

Since the initial reports by Wallen et al. (61) on the hydration of the olefinic bond of oleic acid to 10-hydroxystearic acid by *Pseudomonas* sp., hydration of oleic acid

\[
\text{Oleic acid} + \text{HO} \rightarrow \text{Oleyl alcohol}
\]

By *Euglena gracilis*, *Corynebacterium* sp., and *Rhizopus arrhizus*.

\[
\text{Wax ester (Oleyl oleate)}
\]

*Figure 4* Microbial production of wax esters.
Figure 5  Microbial production of 10-hydroxystearic acid by hydration of oleic acid.

acid catalyzed by Nocardia sp. (62–64), Corynebacterium sp. (65), and Micrococcus sp. (66) have become known.

Anaerobic microbial hydration of cis-9 olefinic bond has been reported to yield 10-hydroxy-12-cis-octadecenoic acid from linoleic acid, 10-hydroxy-12,15-cis,cis-octadecadienoic acid from α-linolenic acid, and 10,12-dihydroxystearic acid from ricinoleic acid as shown in Fig. 6 (67). Recently, hydration catalyzed by Nocardia cholesterolicum has been shown to produce 10-hydroxy-12-cis-octadecenoic acid from linoleic acid in 71% yield and 10-hydroxy-12,15-cis,cis-octadecadienoic acid from α-linolenic acid in 77% yield (68).

A cyanobacterium, Phormidium tenue, converts linoleic acid via 9-hydroperoxy and 13-hydroperoxyoctadecadienoic acids to the corresponding hydroxyoctadecadienoic acids (69). Moderate proportions of 15-hydroxy, 16-hydroxy, and 17-hydroxycis-9-octadecenoic acids have been prepared from oleic acid by Bacillus megaterium and Bacillus pumilus (70).

Strains of Candida tropicalis have been engineered with enhanced ω-hydroxyalase activity to produce ω-hydroxylauric acid from lauric acid as shown in Fig. 7 (71).

A bacterial isolate from Pseudomonas aeruginosa has been found to convert oleic acid to 7,10-dihydroxy-cis-8-octadecenoic acid (Fig. 8), possibly via hydration and hydroxylation (72).

A Flavobacterium sp. has been shown to convert oleic acid to 10-ketostearic acid as the main product (Fig. 9) in addition to 10-hydroxystearic acid (73). Moreover, the above species converts linoleic acid to 10-hydroxy-12-cis-octadecenoic acid (74), and α-linolenic and γ-linolenic acids to 10-hydroxy-12,15-cis,cis-octadecadienoic acid and 10-hydroxy-6,12-cis,cis-octadecadienoic acid, respectively (75). A Staphylococcus sp. has also been found to convert oleic acid to 10-ketostearic acid with 90% conversion and 85% yield (76).
Other examples of microbial biotransformation for the preparation of oxygenated fatty acids include the conversion of oleic acid to 7-hydroxy-17-oxo-9-cis-octadecenoic acid by a *Bacillus* strain (77) and to 10-hydroxy-8-trans-octadecenoic acid (78) and 7,10-dihydroxy-8-trans-octadecenoic acid (79) by *Pseudomonas* sp., formation of 3-R-hydroxy-polyunsaturated fatty acids from fatty acids containing a cis-5,cis-8-diene system by the yeast *Dipodascopsis uninucleata* (80), and the biotransformation of linoleic acid to 8-R-hydroxy-cis-9,cis-12-octadecadienoic acid by the fungus *Leptomitus lacteus* (81). Furthermore, using *Pseudomonas aeruginosa* ricinoleic acid has been converted to 7,10,12-trihydroxy-8-trans-octadecenoic acid (79) via intermediate formation of 10,12-dihydroxy-8-trans-octadecenoic acid (82).
12-Hydroxylauric acid

Figure 7  Microbial production of hydroxy acids by $\omega$-hydroxylation.

3. Polyunsaturated Fatty Acids

Fungi of the Mortierella sp. are capable of converting linoleic acid into $\gamma$-linolenic acid (83) and an oil containing $\alpha$-linolenic acid to an oil containing eicosapentaenoic acid as shown in Fig. 10 (84).

7,10-Dihydroxy-cis-8-octadecenoic acid

Figure 8  Microbial production of dihydroxy acids via hydration/hydroxylation.
4. Dicarboxylic Acids

Species of yeast, *Candida tropicalis*, catalyze the conversion of oleic acid to $\Delta^9$-cis-1,18-octadecenedioic acid (85) (Fig. 11) and elaidic acid to $\Delta^9$-trans-1,18-octadecenedioic acid (86). An industrial strain of *Candida tropicalis* has been engineered to convert methyl myristate, methyl palmitate, methyl stearate, oleic acid and erucic acid to the corresponding $\alpha,\omega$-alkanedicarboxylic acids as outlined in Fig. 11 (71).

A mutant of *Candida tropicalis* efficiently converts linoleic acid into cis-6,cis-9-octadecadienedioic acid, 3-hydroxy-cis-9,cis-12-octadecadienedioic acid, and 3-hydroxy-cis-5,cis-8-octadecadienedioic acid (87). *Candida cloacae* cells oxidize long chain fatty acids to the corresponding dicarboxylic acids (88).

---

**Figure 9**
Microbial production of keto acids.

**Figure 10**
Microbial production of polyunsaturated fatty acids.
Fatty acids and their esters (triacylglycerols) have been converted to methylalkylketones having an alkyl chain one carbon shorter than the substrate using filamentous fungus \textit{Penicillium roquefortii} (89) and the fungus \textit{Trichoderma} sp. (90). A black yeast, \textit{Aureobasidium} sp., produces from lauric acid–rich palm kernel oil \textit{n}-undecane-2-one and \textit{n}-undecane-2-ol in good yields (91). A fungus, \textit{Fusarium avenaceum} sp., produces monohydroxy-\textit{n}-nonane-2-ones from tricaprin (92). These reactions are outlined in Fig. 12.

The fungi \textit{Mucor javanicus} and \textit{Mucor miehei} catalyze the lactonization of \textit{ω}-hydroxy fatty acids, e.g., 15-hydroxypentadecanoic and 16-hydroxyhexadecanoic acids, to the corresponding macrocyclic mono- and oligolactones (93) as shown in Fig. 13.

10-Hydroxystearic acid, prepared by microbial oxidation of oleic acid, has been converted to \textit{γ}-dodecalactone by baker’s yeast, possibly via \textit{β} oxidation (94).

### III. MODIFICATION OF FATS AND OTHER LIPIDS USING ISOLATED ENZYMES

Enzymes of lipid metabolism, isolated from microorganisms, plants, and animal tissues, are highly suitable for biotransformation of fats and other lipids because many of these enzymatic reactions do not require cofactors, and a large number of such enzyme preparations are commercially available. This section will cover the applications of lipolytic enzymes, such as triacylglycerol lipases and phospholipases, as well as the use of oxygenases, lipoxygenases, and epoxide hydrolases in the bio-modification of fats and other lipids.
A. Triacylglycerol Lipases

General properties of triacylglycerol lipases (triacylglycerol acylhydrolases EC 3.1.1.3) are covered in Chapter 26, and the specific applications of triacylglycerol lipases in interesterification reactions are dealt with in Chapters 27 and 28.

Triacylglycerol lipases not only catalyze the hydrolysis; in reaction media with low water content or in the presence of a less polar organic solvent they also catalyze the reverse reaction of lipolysis, i.e., esterification of a fatty acid with an alcohol. Moreover, in media with low water content lipases catalyze a wide variety of interesterification and transesterification reactions, e.g., between triacylglycerols or between a triacylglycerol and a fatty acid (acidolysis) or an alcohol (alcoholysis) or glycerol (glycerolysis) (95–98). The ability of lipases to catalyze the hydrolysis,
esterification, and transesterification of lipids has been exploited in commercial processes for the modification of fats and other lipids.

The substrate specificities and regioselectivities in the hydrolysis of triacylglycerols, catalyzed by some common and commercially available lipases, are summarized in Table 8 (99–103). It is evident that the triacylglycerol lipases have widely varying substrate specificities preferring substrates with long and medium chain fatty acids over the short chain ones and vice versa. Moreover, specificity of lipases for the fatty acids esterified at the sn-1, sn-2, and sn-3 positions of the glycerol backbone vary widely, ranging from nonspecificity for either of the three sn-1, sn-2, and sn-3 positions to strong sn-1,3 or sn-3 specificity.

In general, the substrate specificity and positional specificity of triacylglycerol lipases observed in the hydrolysis reactions are also maintained in the reverse re-

Figure 13  Microbial production of lactones.
Table 8  Specificity of Triacylglycerol Lipases from Different Sources

<table>
<thead>
<tr>
<th>Source of lipase</th>
<th>Fatty acid specificity</th>
<th>Positional specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Microorganisms</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aspergillus niger</td>
<td>S, M, L</td>
<td>sn-1,3 &gt;&gt; sn-2</td>
</tr>
<tr>
<td>Candida antarctica</td>
<td>S &gt; M, L</td>
<td>sn-3</td>
</tr>
<tr>
<td>Candida rugosa (syn. C. cylindracea)</td>
<td>S, L &gt; M</td>
<td>sn-1,2,3</td>
</tr>
<tr>
<td>Chromobacterium viscosum</td>
<td>S, M, L</td>
<td>sn-1,2,3</td>
</tr>
<tr>
<td>Rhizomucor miehei</td>
<td>S &gt; M, L</td>
<td>sn-1,3 &gt;&gt; sn-2</td>
</tr>
<tr>
<td>Penicillium roquefortii</td>
<td>S, M &gt;&gt; L</td>
<td>sn-1,3</td>
</tr>
<tr>
<td>Pseudomonas aeruginosa</td>
<td>S, M, L</td>
<td>sn-1</td>
</tr>
<tr>
<td>Pseudomonas fluorescens</td>
<td>S, L &gt; M</td>
<td>sn-1,2,3</td>
</tr>
<tr>
<td>Rhizopus delmar</td>
<td>S, M, L</td>
<td>sn-1,2,3</td>
</tr>
<tr>
<td>Rhizopus oryzae</td>
<td>M, L &gt; S</td>
<td>sn-1,3 &gt;&gt; sn-2</td>
</tr>
<tr>
<td><strong>Plants</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rapeseed (Brassica napus)</td>
<td>S &gt; M, L</td>
<td>sn-1,3 &gt;&gt; sn-2</td>
</tr>
<tr>
<td>Papaya (Carica papaya) latex</td>
<td>S &gt; M, L</td>
<td>sn-1,3 &gt;&gt; sn-2</td>
</tr>
<tr>
<td><strong>Animal tissues</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Porcine pancreatic</td>
<td>S &gt; M, L</td>
<td>sn-1,3</td>
</tr>
<tr>
<td>Rabbit gastric</td>
<td>S, M, L</td>
<td>sn-3</td>
</tr>
</tbody>
</table>

*S, short chain; M, medium chain; L, long chain.

Superscripts:
- 100. Data from Ref. 100.
- 101. Data from Ref. 101.
- 102. Data from Ref. 102.

Source: Adapted from Ref. 99.

Action of hydrolysis, i.e., esterification (Fig. 14), or in interesterification and transesterification reactions (Fig. 15). The above properties of triacylglycerol lipases permit their use as biocatalyst for the preparation of specific lipid products of definite composition and structure that often cannot be obtained by reactions carried out using chemical catalysts (95). This section outlines some current commercial applications and potentially interesting uses of lipase-catalyzed reactions for the production specialty products from oils and fats.

1. **Structured Triacylglycerols**

a. **Cocoa Butter Substitutes.** Some typical applications of lipase-catalyzed interesterification reactions include the preparation, from inexpensive starting materials, of products resembling cocoa butter in their triacylglycerol structure and physical properties. Commercial processes for the preparation of cocoa butter substitutes involve interesterification of palm oil midfraction with stearic acid or ethyl stearate using sn-1,3-specific lipases, as shown in Fig. 16 (97,104–108).

b. **Human Milk Fat Replacers.** The triacylglycerols of human milk contain the palmitic acid esterified predominantly at the sn-2 position. Structured triacylglycerols resembling triacylglycerols of human milk are produced by transesterification of tripalmitin, derived from palm oil, with oleic acid or polyunsaturated fatty acids, obtained from plant oils, using sn-1,3-specific lipases as biocatalyst as outlined in Fig. 17 (108,109). Such triacylglycerols are used in infant food formulations.

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Figure 14  Specificity of triacylglycerol lipases in hydrolysis and esterification: R₁, R₂, R₃, fatty acids/acyl moieties.

Figure 15  Specificity of triacylglycerol lipases in interesterification and transesterification: R₁, R₂, R₃, fatty acids/acyl moieties.
c. Nutraceuticals. Possible applications of interesterification reactions catalyzed by sn-1,3-specific lipases include, for example, the preparation of structured triacylglycerols for use in specific dietetic products (Nutraceuticals). Thus, interesterification of a common plant oil, such as sunflower oil, with a medium chain fatty acid using a sn-1,3-specific lipase would yield triacylglycerols containing medium chain acyl moieties at the sn-1,3 positions and long chain acyl moieties at the sn-2 position, as shown in Fig. 18 (110–114). Such products, which do not occur in nature and are difficult to prepare by chemical synthesis, may find interesting dietetic applications (115).

Evidence has accumulated lately that nutritional properties of triacylglycerols can be altered beneficially by structuring such lipids, e.g., by inserting certain fatty acyl moieties at specific positions of the glycerol backbone to yield structured triacylglycerols (116–120). Especially, structured triacylglycerols, prepared by lipase-catalyzed transesterification in which the physiologically active ω3 or ω6 polyunsaturated fatty acids, such as docosahexaenoic (DHA) and γ-linolenic (GLA) acids (eicosanoid precursors) are esterified at specific positions of glycerol backbone, as shown in Fig. 19 (121–124) are envisaged to exhibit interesting biological properties (120) that might enable their use in specific nutraceutical products and infant feed (Fig. 19).
Structured triacylglycerols containing two molecules of caprylic acid and one molecule of erucic acid have been prepared by lipase-catalyzed esterification of caprylic acid to monoerucin; the resulting triacylglycerols yield upon subsequent hydrogenation of the erucoyl moieties to behenoyl moieties products resembling Caprenin, a commercially available low-calorie triglyceride (125). Structured triacylglycerols of the type monoglyceride diacetates and diglyceride monoacetates are prepared using \textit{sn}-1,3-specific lipase by transesterification of plant oils with triacetin (126) or an alkyl acetate, e.g., ethyl acetate (127), as shown in Fig. 20.

2. Bioesters, Long Chain Esters, and Flavor Esters

Esterification reactions catalyzed by a nonspecific lipase from \textit{Candida antarctica} (Fig. 21) is being used commercially to produce a wide variety of fatty acid esters, the "bioesters," such as isopropyl myristate, isopropyl palmitate, octyl palmitate, octyl stearate, decyl oleate, and cetyl palmitate, which are used in personal care products (128).

Short chain esters have numerous applications in food industries as flavoring components. Lipase-catalyzed esterification and interesterification for the synthesis of these esters have received considerable attention (129–137).

3. Wax Esters and Steryl Esters

Esterification of mixtures of long chain and very long chain monounsaturated fatty acids with the corresponding mixtures of alcohols using a lipase from \textit{Rhizomucor miehei} (Lipozyme) as catalyst provides wax esters (Fig. 22) in almost theoretical yields (138).

The high rates of interesterification of triacylglycerols with a long chain alcohol (96) indicate that alcoholysis reactions should be useful for the production of wax esters of good commercial value. Using Lipozyme as biocatalyst, wax esters resembling jojoba oil are obtained in high yield by alcoholysis of seed oils from \textit{Sinapis}
alba, Lunaria annua (138), or Crambe abyssinica (139), which contain large proportions of very long chain monounsaturated fatty acids esterified at the sn-1 and sn-3 positions, with very long chain alcohols derived from these oils, as shown in Fig. 23. Similarly, long chain wax esters resembling jojoba oil were obtained in high yields when fatty acids obtained from seed oils of crambe (Crambe abyssinica) and camelina (Camelina sativa) were esterified with oleyl alcohol or the alcohols derived from crambe and camelina oils using Novozym 435 (immobilized lipase B from Candida antarctica) or papaya (Carica papaya) latex lipase as biocatalysts and vacuum was applied to remove the water formed (140). Further examples of lipase-catalyzed preparation of wax esters via esterification (141–146) and interesterification (143–145,147–150) are known.

Unusual wax esters have also been obtained in good yields by lipase-catalyzed reactions, such as esterification of decanol with fatty acids, e.g., 9(10)hydroxymethyl-octadec-10-enoic acid and transesterification of octanol with methyl esters of 9,10-epoxy- or 9-oxodecanoic acids (151).

Lipozyme has been shown to catalyze the esterification of a great variety of carboxylic acids, including short chain, long chain, and branched chain acids to different types of alcohols, ranging from short chain and long chain alkanols to cyclic alcohols (152) giving almost theoretical yields if the water formed by esterification is efficiently removed (105,152).

Esterification catalyzed by immobilized lipases from Rhizomucor miehei (105,152,153) and Candida rugosa (154) as well as surfactant-coated microbial lipases (155) have been carried out for the preparation of a wide variety of alkyl esters of fatty acids in high yields. Moreover, lipase-catalyzed transesterification (alcoholysis) of triacylglycerols with an alcohol, such as n-butanol (156), ethanol or isopropanol (157) provide alkyl esters in high yields, whereby the use of silica gel as an adsorbent for glycerol formed by the reaction greatly enhances the yield (156).

Transesterification (alcoholysis) of low-erucic rapeseed oil with 2-ethyl-1-hexanol, catalyzed by lipase from Candida rugosa, provides 2-ethyl-1-hexyl esters of rapeseed fatty acids in high yields that can serve as a solvent for printing ink (158).

Acid (8:0-14:0) + Isopropanol (Lipase) → (8:0-14:0) Isopropyl ester + Water

**Figure 20** Preparation of structured triacylglycerols of the monoglyceride diacetate and diglyceride monoacetate types by lipase-catalyzed transesterification.

**Figure 21** Lipase-catalyzed synthesis of bioesters.
Butyl esters of fatty acids are useful as lubricants, hydraulic fluids, biodiesel additives, and plasticizers for polyvinylchloride. Butyl oleate has been obtained in high yields by lipase-catalyzed esterification of oleic acid with n-butanol (158).

Long chain alkyl esters of ricinoleic acid have been prepared in high yields by lipase-catalyzed reactions, such as esterification of castor oil fatty acids with a long chain alcohol or transesterification of castor oil triacylglycerols with a long-chain alcohol (159).

Steryl esters of polyunsaturated fatty acids have been obtained by esterification catalyzed by lipase from Pseudomonas sp., but at rather low reaction rates (160). Recently, fatty acyl esters of phytosterols and phytostanols have been obtained at high rates and in near-quantitative yields by esterification of the sterols with the fatty acids or their transesterification with alkyl esters of fatty acids, both under vacuum using Candida rugosa lipase as biocatalyst (161). Such steryl esters are being used recently as blood cholesterol–lowering food supplements added to margarines (162).

4. Monoacylglycerols and Diacylglycerols

Lipase-catalyzed partial hydrolysis of oils (163,164) and esterification of fatty acids with glycerol (165–174) have been carried out for the production of monoacylglycerols (Fig. 24). Lipase-catalyzed esterification of glycerol with fatty acids under vacuum provides symmetrical 1,3-diacylglycerols in good yields (175). Moreover, lipase-catalyzed transesterification of glycerol with an alkyl ester of a fatty acid (170,171) or of triacylglycerols with an alcohol, such as ethanol (176) or n-butanol (177), provides good yields of monoacylglycerols.

Interesterification (glycerolysis) of triacylglycerols with glyceroglycerol, catalyzed by lipases, as shown in Fig. 25, has been by far most successful for the preparation of monoacylglycerols (178–187). Diacylglycerols have also been prepared in high yields by glycerolysis of hydrogenated beef tallow, catalyzed by lipase from Pseudomonas sp. (188).
Monoacylglycerols of less common fatty acids, such as 9(10)acetonoyloctadecanoic acid, have been obtained in high yields by one-pot reaction of this acid with glycerol and lipase in the presence of phenylboronic acid as a solubilizing agent (151).

5. Lactones and Estolides

Musk lactones are used in the fragrance industry. Hexadecanolide, a musk monolactone, has been obtained in good yields by intramolecular lactonization of 16-hydroxyhexadecanoic acid, catalyzed by immobilized lipase from *Candida antarctica*, whereby oligolactones are not formed by intermolecular lactonization (189).

Reaction of lesquerolic (14-hydroxy-11-eicosenoic) acid with oleic acid, catalyzed by lipase from *Candida rugosa*, produces mainly monoestolides containing one molecule each of the hydroxy acid and oleic acid per molecule, whereas the corresponding reaction, catalyzed by *Pseudomonas* sp. lipase, produces substantial proportions of monoestolides containing two molecules of lesquerolic acid per molecule besides diestolides (190). Properties of mono- and polyestolides, synthesized chemically, can be substantially improved by esterification of the estolides with fatty alcohols or \( \alpha,\omega \)-diols, catalyzed by lipase from *Rhizomucor miehei* (190).

**Figure 24** Preparation of monoacylglycerols by lipase-catalyzed partial hydrolysis of fats (top) and esterification of fatty acids with glycerol (bottom).

**Figure 25** Preparation of mono- and diacylglycerols by lipase-catalyzed interesterification of fats with glycerol (glycerolysis).
6. Fatty Acid Esters of Sugars, Alkylglycosides, and Other Hydroxy Compounds

Lipase-catalyzed esterification has been carried out for the synthesis of fatty acyl esters of carbohydrates that can be used as emulsifiers. Esters of monosaccharides and disaccharides (191–202) as well as those of sugar alcohols (192,193,195, 196,203) and other polyols (204) have been prepared in good yields and excellent regioselectivity by esterification catalyzed by microbial lipases (Fig. 26). Transesterification of sugars with short chain alkyl esters of fatty acids also provide sugar esters in good yields (191,205).

Recently, lipase-catalyzed esterification and transesterification reactions using less toxic solvents, such as tert-butanol (206,207) or acetone (208,209), have been used successfully for the preparation of sugar esters.

Triacylglycerols, contained in common fats and oils, as well as wax esters of jojoba oil, have been transesterified with various sugar alcohols in pyridine using lipases to yield primary monoesters of sugar alcohols having excellent surfactant properties (210). Transesterification (211,212) and esterification (213,214) reactions, catalyzed by lipases, have also been applied for the preparation of fatty acid esters of alkyl glycosides (Fig. 27).

Figure 26 Preparation of sugar esters of fatty acids by lipase-catalyzed esterification.

Figure 27 Preparation of esters of alkylglycosides by lipase-catalyzed esterification.
Fatty acid esters of polyols are useful as surfactants. Polyglycerol–fatty acid esters have been prepared in good yields by transesterification of polyglycerol, adsorbed on silica gel, with methyl esters of fatty acids (215).

Propylene glycol monoesters, suitable as emulsifiers, have been prepared in good yields by reacting a fatty acid anhydride with 1,2-propanediol in the presence of lipase from a *Pseudomonas* sp. (216). Esterification of eicosapentaenoic and docosahexaenoic acids with 1,2-propanediol in the presence of lipase from *Rhizomucor miehei* provides propylene glycol monoester emulsifiers that are potentially beneficial to health (217).

Polyethylene glycol esters of fatty acids, widely used as nonionic surfactants, have been prepared in essentially quantitative yields by esterification of oleic acid with polyethylene glycol, catalyzed by lipase from *Rhizomucor miehei* (218).

Medium and long chain alcohols (C₈–C₁₆) have been efficiently esterified with lactic acid and glycolic acid using lipase B from *Candida antarctica* (Novozym 435) as biocatalyst (219). Using the same lipase ethyl lactate has been transesterified with β-n-octyl-D-glucopyranoside to obtain β-n-octyl-D-glucopyranosyl lactate in high yield (220).

Ascorbyl palmitate, used as antioxidant in foods and cosmetics, has been prepared in good yields by esterification of ascorbic acid with palmitic acid using lipase from *Bacillus stearothermophilus* SB 1 (221) and *Candida antarctica* (Novozym 435) (222). Esterification of cinnamic acid with 1-octanol using Novozym 435 also provides the octyl ester in moderate yields (222). Similarly, using lipase B from *Candida antarctica* (Chirazyme L2) 6-O-palmitoyl-L-ascorbic acid and 6-O-eicosapentaenoyl-L-ascorbic acid have been prepared, respectively, via transesterification with vinyl palmitate (223) and condensation with eicosapentaenoic acid (224). Transesterification of L-methyl lactate with ascorbic acid or retinol using Novozym 435 gives high yields of ascorbyl-L-lactate and retinyl-L-lactate, respectively (225).

### 7. Amides

Reaction of a triacylglycerol mixture, such as soybean oil, with lysine, catalyzed by lipase from *Rhizomucor miehei* yields acyl amides, i.e., N-acyllysines (226). Reaction of ethyl octanoate with ammonia (ammonolysis), catalyzed by lipase from *Candida antarctica*, provides octanamide in near-quantitative yields (227). One pot enzymatic synthesis of octanamide in high yield via esterification of octanoic acid with ethanol, followed by ammonolysis of the resulting ethyl octanoate, both reactions being conducted using the lipase from *Candida antarctica* has also been reported (227). Lipase-catalyzed direct amidation of carboxylic acids by ammonia and ammonium salts has been reported (228) and various applications of lipase-catalyzed aminolysis and ammonolysis have been recently reviewed (229).

### 8. Fatty Acids

Lipase-catalyzed hydrolysis can be applied for the production of fatty acids from fats using, e.g., a nonspecific lipase preparation from *Candida rugosa* (syn. *Candida cylindracea*) (230–238), *Pseudomonas* sp. (239–243), *Aspergillus* sp. (244), *Thermomyces lanuginosus* (245), and *Chromobacterium viscosum* (246) as alternative mild processes compared to drastic “steam splitting” (Fig. 28). Specifically, fatty acids have been obtained by lipase-catalyzed hydrolysis of technically important fats, such as animal fats (236), castor oil (243), palm stearin (242), and the thermally
labile high-\(\alpha\)-linolenic acid–containing perilla oil (238). More recently, enzymatic “presplitting” of oils prior to steam splitting is being considered as an economically feasible alternative to complete enzymatic hydrolysis or steam splitting alone (247).

a. Fatty Acid Concentrates. Triacylglycerol lipases from various organisms have one common feature in their selectivity toward groups of fatty acids/acyl moieties having olefinic bonds at definite positions (249) or geometric configuration (249). Thus, several lipases from microorganisms, plants and animal tissues discriminate against fatty acids/acyl moieties having a \(cis\)-4, \(cis\)-6, or a \(cis\)-8 double bond as substrates in hydrolysis, esterification, and interesterification reactions, as summarized in Table 9.

Data presented in Fig. 29 show for example the substrate specificity in esterification of a wide variety of fatty acids with \(n\)-butanol using the latex from papaya plant (\(Carica papaya\)) as biocatalyst (256). In these studies a mixture of the fatty acid examined and the reference standard, myristic acid, at equal molar concentrations in \(n\)-hexane was reacted with \(n\)-butanol using the above biocatalyst and the course of formation of butyl esters under competitive conditions was followed. The competitive factor \(\alpha\) was determined according to Rangheard et al. (252) from the concentrations of the two substrates (\(Ac1X\) and \(Ac2X\)) at time \(X\) by the equation:

\[ \alpha = \frac{V_{Ac1X}/K_{Ac1X}}{V_{Ac2X}/K_{Ac2X}} \]

where \(V\) is maximal velocity and \(K\) is the Michaelis constant. The competitive factor

![Figure 28](image.png)

**Figure 28** Preparation of fatty acids and glycerol by lipase-catalyzed hydrolysis of fats.

<table>
<thead>
<tr>
<th>Table 9</th>
<th>Specificity of Triacylglycerol Lipases from Different Sources Toward Various Fatty Acids/Acyl Moieties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Source of lipase</td>
<td>Discrimination against*</td>
</tr>
<tr>
<td>Microorganisms</td>
<td></td>
</tr>
<tr>
<td><em>Candida rugosa</em> (syn. <em>C. cylindracea</em>)</td>
<td>all-(cis)-4,7,10,13,16,19-DHA, petroselinic acid, GLA (all-(cis)-6,9,12-octadecadienoic) acid, stearidonic acid, dihomoy-(\gamma)-linolenic acid (all-(cis)-6,9,12-eicosatrienoic) acid</td>
</tr>
<tr>
<td><em>Penicillium cyclopium</em></td>
<td></td>
</tr>
<tr>
<td><em>Penicillium</em> sp. (lipase G)</td>
<td></td>
</tr>
<tr>
<td><em>Rhizomucor miehei</em></td>
<td></td>
</tr>
<tr>
<td><em>Rhizopus arrhizus</em></td>
<td></td>
</tr>
<tr>
<td>Plants</td>
<td></td>
</tr>
<tr>
<td>Rape (<em>Brassica napus</em>) seedlings</td>
<td>DHA, petroselinic acid, GLA, stearidonic, and dihomoy-(\gamma)-linolenic acids</td>
</tr>
<tr>
<td>Papaya (<em>Carica papaya</em>) latex</td>
<td></td>
</tr>
<tr>
<td>Animal tissues</td>
<td></td>
</tr>
<tr>
<td>Porcine pancreas</td>
<td>DHA, petroselinic acid, GLA, and stearidonic acid</td>
</tr>
</tbody>
</table>

*\(DHA\), docosahexaenoic acid; GLA, \(\gamma\)-linolenic acid.*
Specificity constants in the esterification of mixtures of myristic acid (reference standard) and individual fatty acids with n-butanol in hexane using papaya latex as biocatalyst. (From Ref. 256.)

\[ \alpha = \log\left(\frac{Ac1X0}{Ac1X}\right) / \log\left(\frac{Ac2X0}{Ac2X}\right) \]

From the competitive factor specificity constant was calculated as 1/\(\alpha\) with reference to the specificity constant of myristic acid taken as 1.00. The higher the specificity constant of a fatty acid the greater is the specificity of the biocatalyst for that particular fatty acid. The data presented in Fig. 29 show that the fatty acids having a cis-4, cis-6, or a cis-8 double bond are poor substrates in esterification reactions as compared to those having a cis-5 or cis-9 double bonds or fatty acids having hydroxy, epoxy, or cyclopentenyl groups.

The above substrate specificities have been utilized for the enrichment of definite fatty acids or their derivatives from mixtures via kinetic resolution (257,258), e.g., by selective hydrolysis as shown in Scheme 3 and the following examples.

Despite relatively high prices of lipase preparations, lipase-catalyzed hydrolysis could be economically attractive for the preparation of specific products of high commercial value, such as polyunsaturated (\(\omega3\)) fatty acid concentrates via selective hydrolysis of marine oils, catalyzed by fatty acid-specific lipases that enable the enrichment of docosahexaenoic 22:6 \(\omega3\) and eicosapentaenoic 20:5 \(\omega3\) acids in the unhydrolyzed acylglycerols, as outlined in Fig. 30 (257,259–263). Such polyunsaturated fatty acids which are interesting as dietetic products (264) cannot be obtained by conventional steam splitting without substantial decomposition.
Scheme 3  Principle of kinetic resolution via lipase-catalyzed hydrolysis.

In a commercial process fish oil is partially hydrolyzed by Candida rugosa lipase to yield an acylglycerol fraction enriched in 20:5 ω3, and especially in 22:6 ω3; the acylglycerol fraction is subsequently isolated by evaporation and converted to triacylglycerols via hydrolysis and reesterification, both catalyzed by R. miehei lipase (265). Using Rhizopus delemar lipase, selective esterification of tuna oil fatty acids with lauryl alcohol, extraction of the unreacted fatty acids and their repeated esterification with lauril alcohol has resulted in an unesterified fatty acid fraction containing 91% 22:6 ω3 (266). Selective interesterification of tuna oil triacylglycerols with ethanol using Rhizomucor miehei lipase as biocatalyst yields an acylglycerol fraction containing 49% 22:6 ω3, whereas selective esterification of tuna oil fatty acids with ethanol yields an unesterified fatty acid fraction containing 74% 22:6 ω3 (267).

Fatty acids generated by lipase-catalyzed hydrolysis of a commercial single-cell oil from Mortierella alpina have been subjected to selective esterification with lauryl alcohol, catalyzed by lipase from Candida rugosa. This leads to an increase in the arachidonic acid content from 25% in the starting fatty acid mixture to over 50% in the fatty acids that remained unesterified (268).

Also γ-linolenic acid, a constituent of certain seed oils, such as borage oil and evening primrose oil, can be prepared as a concentrate together with linoleic acid by lipase-catalyzed selective hydrolysis (Fig. 31) under mild conditions and typical data obtained with lipase from Candida rugosa are shown in Fig. 32 (269).

Figure 30  Preparation of concentrates of docosahexaenoic acid (DHA) via lipase-catalyzed selective hydrolysis of marine oils.
Figure 31 Preparation of concentrates of γ-linolenic acid (GLA) via lipase-catalyzed selective hydrolysis of borage oil or evening primrose oil.

Very recently, lipase-catalyzed selective hydrolysis of a microbial oil from Mortierella alpina has been employed to enrich arachidonic acid in the acylglycerols (270).

The ability of lipase preparations from plants and microorganisms to discriminate against fatty acids/acyl moieties having cis-4-, cis-6, or cis-8 double bonds (Table 9) has been utilized for the enrichment of γ-linolenic acid from fatty acid mixtures, derived from plant and microbial oils, by selective esterification of the fatty acids, other than γ-linolenic acid, with n-butanol as outlined in Fig. 33 (271–276). Similarly, lipase-catalyzed esterification has been applied to enrich docosahexaenoic acid from fatty acid mixtures, derived from marine oils (272,276). Such concentrates might find nutraceutical applications in capsules.

Typical data on enrichment of GLA via lipase-catalyzed selective esterification of fatty acids from borage oil with n-butanol are given in Fig. 34.

Conjugated linoleic acids (CLA), predominantly a mixture of cis-9,trans-11-octadecadienoic and trans-10,cis-12-octadecadienoic acids, have gained some interest as beneficial supplements for foods and feeds due to their potentially anticarcinogenic and immunological properties. However, it is not known which of the two major isomers is physiologically more active. With this background, a mixture con-
Figure 33 Preparation of concentrates of γ-linolenic acid (GLA) via lipase-catalyzed selective esterification of fatty acids from borage oil or evening primrose oil with n-butanol.

taining the two CLA isomers in equal amounts was subjected to esterification with dodecanol using lipase from Geotrichum candidum as catalyst (277). This resulted in selective esterification of the cis-9,trans-11 isomer and enrichment of the trans-10,cis-12 isomer in the unesterified fatty acid fraction. Separation of the two fractions by molecular distillation yielded an ester fraction containing 91% cis-9,trans-11 isomer and a fatty acid fraction containing 82% trans-10,cis-12 isomer.

Selective hydrolysis of high-erucic oils, catalyzed by lipases from Geotrichum candidum (278,279) and Candida rugosa (279,280), leads to enrichment of erucic acid in the unhydrolyzed acylglycerols, as outlined in Fig. 35. The diacylglycerols formed by hydrolysis using the lipase from Candida rugosa contain as much as 95% of erucic acid (279). In the esterification of individual fatty acids with n-butanol, catalyzed by lipase from Geotrichum candidum, erucic acid is discriminated against (281,282).

The selectivity of commercial lipases towards very long chain monounsaturated fatty acids (VLCMFAs) in hydrolysis and transesterification reactions has been determined using high-erucic oils from white mustard (Sinapis alba), oriental mustard (Brassica juncea), and honesty (Lunaria annua) seeds (127). The lipases from Candida rugosa and Geotrichum candidum selectively cleave the C18 fatty acids from

![Diagram](image-url)

Figure 34 Enrichment of γ-linolenic acid (GLA) from borage oil fatty acids by selective esterification with n-butanol, catalyzed by triacylglycerol lipase from Rhizomucor miehei (Lipozyme) according to Ref. 275: reaction temperature, 60°C; reaction time, 2 hours; degree of esterification, 91%.
the triacylglycerols, which results in enrichment of these fatty acids from about 40% in the starting oil to approximately 60% and 89%, respectively, in the fatty acid fraction; concomitantly, the level of erucic acid and the other VLCMFA is raised in the acylglycerol fraction from 51% in the starting oil to about 80% and 72%, respectively, as shown in Fig. 36.

The \( sn \)-1,3-specific lipases from porcine pancreas, \textit{Chromobacterium viscosum}, \textit{Rhizopus arrhizus}, and \textit{Rhizomucor miehei} selectively cleave the VLCMFA, esterified almost exclusively at the \( sn \)-1,3 positions of the high-erucic triacylglycerols, which results in enrichment of the VLCMFA in the fatty acid fraction to 65–75%, whereas the C\textsubscript{18} fatty acids are enriched in the acylglycerol fraction (127).

Selective hydrolysis of erucic acid– and nervonic acid–rich triacylglycerols of \textit{Lunaria annua}, catalyzed by the lipase from \textit{Candida rugosa}, leads to preferential cleavage of the C\textsubscript{18} fatty acids, resulting in their enrichment from 36% in the starting oil to 79% in the fatty acids; concomitantly, the VLCMFA are enriched in the di- and triacylglycerols. The diacylglycerols, the major (55%) products of lipolysis, are almost exclusively (>99%) composed of VLCMFA (127).

\textbf{Figure 35} Enrichment of very long chain monounsaturated fatty acids from high erucic acid seed oils via lipase-catalyzed selective hydrolysis.

\textbf{Figure 36} Enrichment of very long chain monounsaturated fatty acids (VLCMFA = eicosenoic + erucic + nervonic) from white mustard seed oil by selective hydrolysis catalyzed by triacylglycerol lipase from \textit{Candida cylindracea} (syn. \textit{C. rugosa}) according to Ref. 127: reaction temperature, 20°C; reaction time, 1.25 hours; degree of hydrolysis, 49%.
Lipase-catalyzed selective hydrolysis of triacylglycerols from meadowfoam oil or selective esterification of meadowfoam fatty acids have been reported for the enrichment of cis-5-eicosenoic acid in the acylglycerols and unesterified fatty acids, respectively (283).

Fennel (*Foeniculum vulgare*) oil has been selectively hydrolyzed using a lipase from *Rhizopus arrhizus* for the enrichment of petroselinic acid in the unhydrolyzed acylglycerols (284). The acylglycerols were separated from the fatty acids using an ion exchange resin and subsequently hydrolyzed by the lipase from *Candida rugosa* to yield a fatty acid concentrate containing 96% of petroselinic acid.

Highly purified concentrate of petroselinic acid has also been prepared from fatty acids of coriander oil via selective esterification with n-butanol, catalyzed by lipase from germinating rapeseed (Fig. 37) (276).

Fatty acid concentrates containing 85% hydroxy acids, such as lesquerolic (14-hydroxy-cis-11-eicosenoic) and auricolic (14-hydroxy-cis-11-cis-17-eicosadienoic) acids, have been prepared from lesquerella oil by their selective cleavage catalyzed by lipase from *Rhizopus arrhizus* (285).

Fatty acids of *Biota orientalis* seed oil have been selectively esterified with n-butanol using lipase from *Candida rugosa* to enrich cis-5-polyunsaturated fatty acids, e.g., all-cis-5,11,14-octadecatrienoic and all-cis-5,11,14,17-octadecatetraenoic acid; the level of total cis-5-polyunsaturated fatty acids is raised from about 15% in the starting material to about 73% in the unesterified fatty acids (286). Selective hydrolysis of the seed oil of *Biota orientalis* by the lipase from *Candida rugosa* leads to enrichment of the cis-5-polyunsaturated fatty acids in the acylglycerols to about 41% (286). Apparently, fatty acids/acyl moieties having a cis-5 double bond are also discriminated against by some lipases.

A recent comprehensive review (248) covers the applications of lipase-catalyzed reactions for the enrichment of fatty acids via kinetic resolution.

### 9. Phospholipids

Interestingly, not only phospholipases A1 and A2, as described in Sec. B, but also some triacylglycerol lipases cleave the fatty acids from the sn-1 and/or sn-2 positions of diacylglycerophospholipids and also catalyze ester exchange reactions to modify the composition of the acyl moieties at the sn-1 and/or sn-2 positions of glycerophospholipids. Interestingly, triacylglycerol lipases, such as those from *Rhizopus ar- rhizus* and *Rhizomucor miehei*, have recently been found to be also able to catalyze the acyl exchange of galactolipids, e.g., via acidolysis of heptadecanoic acid with digalactosyldiacylglycerols (DGDG) (287). Modification of phospholipids using triacylglycerol lipases will be covered in Sec. B.1.

**Figure 37** Preparation of concentrates of petroselinic acid via lipase-catalyzed selective esterification of fatty acids from coriander oil with n-butanol.
B. Phospholipases

Figure 38 shows the reactions catalyzed by phospholipases (see also Chapter 26). Phospholipases A₁ and A₂ hydrolyze the acyl moieties from the \( sn-1 \) and \( sn-2 \) positions, respectively, of glycerophospholipids, such as diacylglycerophosphocholines. Phospholipase C cleaves the polar head groups, such as phosphocholine or phosphoethanolamine residues, esterified at the \( sn-3 \) position of these phospholipids yielding diacylglycerols. Phospholipase D cleaves the bases or alcohols, such as choline or ethanolamine, from these phospholipids yielding phosphatidic acids. Under certain conditions, such as in the presence of less polar organic solvents at low water content, most of the above reactions can be reversed to modify the composition of the acyl moieties or the head groups of the phospholipids.

Phospholipids, such as commercial “soya lecithin” or “egg lecithin,” are widely used for their emulsifying and other functional properties in food, cosmetic, and pharmaceutical products. The composition of phospholipids can be altered to modify their properties by chemical reactions (288) or enzymatic reactions catalyzed by phospholipases (95,118,289).

1. Phospholipids Modified by Phospholipases A₁ and A₂ and Triacylglycerol Lipases

The following examples show the various possibilities of modifying the composition of acyl moieties of phospholipids by interesterification reactions catalyzed by phospholipase A₁, phospholipase A₂, or triacylglycerol lipase. Such modified phospholipids may find interesting biomedical applications.

Selective hydrolysis of diacylglycerophospholipids, catalyzed by phospholipase A₂ or A₁, yields 1-acyl- or 2-acyllysoglycerophospholipids, respectively (118) (Fig. 38). Phospholipases A₁ and A₂ as well as regiospecific or nonregiospecific triacylglycerol lipases have been found to cleave the fatty acids from the \( sn-1 \) and/or \( sn-2 \) positions of diacylglycerophospholipids to yield \( sn-1- \) or \( sn-2- \) lysoglycerophospholipids with interesting functional properties (290–295). Studies on hydrolysis of soybean phospholipids have revealed that fungal triacylglycerol lipase preparations that also contain phospholipase A₁ and A₂ activities as well as lysophospholipase activity are more efficient in the cleavage of fatty acids than fungal and mammalian enzyme preparations that have only phospholipase A₁ and/or A₂ activities (296). Moreover, fungal preparations of both triacylglycerol lipases as well as phospholipase A₁ cleave in the course of time the fatty acids esterified at both \( sn-1 \) and \( sn-2 \) positions of diacylglycerophospholipids yielding completely deacylated products, e.g., glycerylphosphorylcholine (296). However, phospholipase A₂ preparations of fungal as well as mammalian pancreatic origin yield primarily \( sn-1- \)-acyllysophospholipid by selective partial deacylation at the \( sn-2 \) position (296).

Esterification of \( sn-1 \)-acyllysoglycerophosphocholines with eicosapentaenoic acid and docosahexaenoic acid, catalyzed by porcine pancreatic phospholipase A₂ in a microemulsion system containing small amounts of water, has been carried out to prepare diacylglycerophosphocholines containing well over 30% \( \omega-3 \) polyunsaturated fatty acids (\( \omega-3 \) polyunsaturated fatty acids, PUFAs) as outlined in Fig. 39 (297). Transesterification of phosphatidylcholine with ethyl eicosapentaenoate, catalyzed by porcine pancreatic phospholipase A₂ in the presence of toluene, has led to about 14% incorporation of eicosapentaenoyl moieties into phosphatidylcholine (298).
Figure 38  Reactions catalyzed by phospholipases (PL) A₁, A₂, C, and D: R₁, R₂, fatty acids/acyl moieties; X, base or alcohol (e.g., choline, ethanolamine, etc.).

Long chain PUFAs from fish oil, dissolved in propane (299) or isooctane (300) have been esterified to sn-1-acyl-lysophosphatidylcholine to an extent of about 20 to 25% using porcine pancreatic phospholipase A₂ as biocatalyst. Similarly, phospholipase A₂–mediated esterification of eicosapentaenoic acid to lysophosphatidylcholine in the presence of formamide has been used to prepare therapeutic phospholipids in yields of about 60% (301).

Figure 39  Preparation of structured phospholipids by esterification of lysophospholipids with ω3 polyunsaturated fatty acids (ω3 PUFA) catalyzed by phospholipase A₂.
Lysophosphatidic acid has been prepared in a yield of 32\% by direct solvent-free esterification of fatty acids to sn-glycerol-3-phosphate, catalyzed by triacylglycerol lipase from \textit{Rhizomucor miehei} (302). Immobilized \textit{sn}-1,3-specific triacylglycerol lipase from \textit{Rhizopus arrhizus} has been found to efficiently catalyze the transesterification of \textit{dl}-glycerol-3-phosphate with lauric acid vinyl ester yielding lysophosphatidic acids (1-acyl-\textit{rac}-glycerol-3-phosphate) and phosphatidic acids (1,2-diacyl-\textit{rac}-glycerol-3-phosphate) in a total conversion of >95\% (303). The conversions were lower (55\%) with oleic acid as acyl donor in the corresponding esterification reaction (303).

Transesterification of \textit{l}-\textit{\alpha}\textsuperscript{-glycerophosphocholine} with vinyl esters of fatty acids such as vinyl laurate, catalyzed by \textit{Candida antarctica} lipase B (Novozym 435) in the presence of \textit{tern}-butanol, gives predominantly 1-acyllysophosphatidylcholine with high (>95\%) conversion (304). Similarly, esterification of fatty acids with \textit{l}-\textit{\alpha}\textsuperscript{-glycerophosphocholine}, catalyzed by \textit{Rhizomucor miehei} lipase (Lipozyme IM) in the presence of dimethylformamide, produces 1-acyllysophosphatidylcholine with high (90\%) conversion (305).

Transesterification reactions, such as \textit{acidolysis}, i.e., exchange of the constituent fatty acids of diacylglycerophosphocholines, have been carried out against other fatty acids added as reaction partners using \textit{sn}-1,3-specific triacylglycerol lipase from \textit{Rhizopus delemar} as biocatalyst (306) (Fig. 38). Similarly, transesterification of diacylglycerophospholipids, catalyzed by \textit{sn}-1,3-specific or nonspecific triacylglycerol lipases, has been applied to modify the fatty acid composition of diacylglycerophospholipids, specifically at the \textit{sn}-1 position or at both \textit{sn}-1 and \textit{sn}-2 positions (295,296,307–312). In transesterification of phosphatidylcholine with a fatty acid, catalyzed by triacylglycerol lipases from \textit{Rhizopus delemar}, \textit{Rhizomucor miehei} (308), or \textit{Rhizopus arrhizus} (309,310) acyl exchange occurs almost exclusively at the \textit{sn}-1 position. Polysaturated fatty acids, especially \textit{\omega}-PUFA, have been incorporated into phospholipids by transesterification catalyzed by \textit{sn}-1,3-specific triacylglycerol lipases from \textit{Rhizopus delemar} (311) and \textit{Rhizomucor miehei} (312), as outlined in Fig. 40.

Transesterification of eicosapentaenoic acid with phosphatidylcholine from soybean, catalyzed by an \textit{sn}-1,3-specific triacylglycerol lipase from \textit{Rhizomucor miehei} in the presence of a combination of water and propylene glycol yields a therapeutically beneficial phospholipid (301).

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure40.png}
\caption{Preparation of structured phospholipids by transesterification of diacylglycerophospholipids with \textit{\omega}-PUFA polyunsaturated fatty acids catalyzed by triacylglycerol lipases.}
\end{figure}
Moreover, transesterification, such as alcoholyis of a phospholipid with an alcohol, such as ethanol, isopropanol, or n-butanol, catalyzed by an sn-1,3-specific lipase from *Rhizomucor miehei*, has been employed for the preparation of lysophospholipids (Fig. 41) (313).

2. Phospholipids Modified by Phospholipase D

The following examples show the various possibilities of modifying the composition of polar head groups of phospholipids by phospholipase D-catalyzed reactions. Such modified phospholipids may find interesting biomedical applications. 

*Transphosphatidylation* (base exchange) reactions of phospholipids, catalyzed by phospholipase D (Fig. 38), can be utilized for the preparation of specific phospholipids. For example, phospholipase D-catalyzed transphosphatidylation reaction of egg lecithin (predominantly phosphatidylcholines and phosphatidylethanolamines) with glycerol yields phosphatidylylycerols with the simultaneous formation of choline and ethanolamine (Fig. 42) (314,315).

Phosphatidylglycerols may find biomedical applications as physiologically active pulmonary surfactant (316). Efficient methods have been described for the preparation of phosphatidylglycerols from phosphatidylcholines and glycerol by transphosphatidylation catalyzed by phospholipase D (317–319). Transphosphatidylation reactions catalyzed by phospholipase D have also been carried out to convert ethanolamine plasmalogens to their dimethyllethanolamine or choline analogs (320) and to obtain phosphatidylethanolamines (321) or phosphatidylserines (322,323) from phosphatidylcholines (Fig. 43).

Phosphatidylcholine content of commercial lecithins has been increased by transphosphatidylation of lecithins with choline chloride catalyzed by phospholipase D (324).

Another example of phospholipase D-catalyzed transphosphatidylation reaction is the synthesis of structural analogs of platelet-activating factor, PAF (1-O-alkyl-2-acetyl-sn-glycero-3-phosphocholine) by the replacement of choline by primary cyclic alcohols (325). Moreover, transphosphatidylation of pure glycerophospholipids or commercial products, e.g., soy lecithin or egg lecithin, with a sugar, such as glucose, using phospholipase D from an *Actinomadura* sp. as biocatalyst and diethyl ether or tert-butanol as solvent affords the phosphatidylglucose or other phosphatidylsaccharides in yields as high as 85% (Fig. 44) (326).

Transphosphatidylation of phosphatidylcholine with 1-monolauroyl-rac-glycerol, catalyzed by phospholipase D from *Streptomycetes* sp. yields 1-lauroyl-phosphatidylglycerol which has been subsequently cleaved by phospholipase C from *Bacillus cereus* to yield 1-lauroyl-rac-glycerophosphate (327). Similarly, transphosphatidylation of phosphatidylcholine with 1-lauroyl-dihydroxyacetone, catalyzed by phospholipase D, yields 1-lauroyl-phosphatidylhydroxyacetone, which has been subsequently cleaved by phospholipase C to yield 1-lauroyl-dihydroxyacetonephosphate (327).

C. Other Enzymes

1. Lipooxygenases

Lipooxygenase from soybean converts linoleic acid or other compounds having a cis,cis-1,4-pentadiene system to conjugated hydroperoxides (Fig. 45) (328). Soybean
**Figure 41** Transesterification of phosphatidylcholines with an alcohol (alcoholysis) catalyzed by triacylglycerol lipases for the preparation of lysophosphatidylcholines.

\[
\text{SN-1,3-Specific triacylglycerol lipase} \\
\text{Phosphatidylcholine + Alcohol} \quad \xrightarrow{\text{Lipase}} \quad \text{Lysophosphatidylcholine + Alkyl esters} 
\]

**Figure 42** Transphosphatidylation of phosphatidylcholines with glycerol catalyzed by phospholipase D for the preparation of phosphatidylglycerols.

\[
\text{Phospholipase D} \\
\text{Phosphatidylcholine + Glycerol} \quad \xrightarrow{\text{Phospholipase D}} \quad \text{Phosphatidylglycerol + Choline} 
\]

**Figure 43** Transphosphatidylation reactions catalyzed by phospholipase D.

**Figure 44** Transphosphatidylation of phosphatidylcholines with glucose catalyzed by phospholipase D for the preparation of phosphatidylglucose.

\[
\text{Phospholipase D} \\
\text{Phosphatidylcholine + Glucose} \quad \xrightarrow{\text{Phospholipase D}} \quad \text{Phosphatidylglucose + Choline} 
\]

**Figure 45** Preparation of hydroperoxides from linoleic acid by soybean lipoxygenase.

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Lipoxygenase has been used for example to prepare 9-hydroperoxy-γ-linolenic acid from γ-linolenic acid (329), hydroperoxides of acylglycerols and phospholipids (330,331) and dimers of linoleic acid (332).

Recently, lipoxygenase immobilized in a packed-bed column reactor has been used to convert linoleic acid to hydroperoxyoctadecadienoic acid (333). Furthermore, a double-fed batch reactor fed with a mixture of linoleic acid together with a crude lipoxygenase extract from defatted soybean flour has been used to obtain hydroperoxyoctadecadienoic acid, which is subsequently reduced in situ by cysteine, also contained in the reactor, to give 13(S)-hydroxy-9-cis,12-cis-octadecadienoic acid in high yield (334).

2. Oxygenases

Crude enzyme preparations from a variety of organisms have been shown to exhibit interesting activities that can be utilized for the biotransformation of fats and other lipids. For example, enzyme preparation from plants have been shown to catalyze ω-hydroxylation (Fig. 46) (335) and epoxidation (Fig. 47) (336) of fatty acids. Recently, a peroxygenase isolated from oat (Avena sativa) has been immobilized on synthetic membranes and employed for the epoxidation of oleic acid using hydrogen peroxide or organic hydroperoxides as oxidants (337).

Enzyme preparations containing alcohol oxidase, isolated from the yeast Candida tropicalis, catalyze the oxidation of long chain alcohols, diols, and ω-hydroxy fatty acids to the corresponding aldehydes (Fig. 48) (338). Alcohol oxidase preparations, isolated from the yeast Candida maltosa, catalyze the oxidation of 1-alkanols and 2-alkanols to the corresponding aldehydes and ketones, respectively (339).

3. Epoxide Hydrolases

Epoxide hydrolases (EC 3.3.2.3) are ubiquitous in nature (340). They catalyze the hydrolysis of epoxides to vicinal diols. In particular, epoxide hydrolases from higher plants are well characterized (341–345).

Epoxide hydrolase from soybean seedlings catalyzes the hydration of cis-9,10-epoxystearic acid to threo-9,10-dihydroxystearic acid (Fig. 49) (341):

The two positional isomers of linoleic acid monooepoxides are hydrated to their corresponding vic-diols by soybean fatty acid epoxide hydrolase (Fig. 50) (342).

The epoxide hydrolase has been found to be highly enantioselective with strong preference for the enantiomers cis-9R,10S-epoxy-cis-12-octadecenoic acid and cis-12R-,13S-epoxy-cis-9-octadecenoic acid, respectively (342–344).

Strong enantioselection has also been reported for rabbit liver microsomal epoxide hydrolase (346). Cloning and expression of soluble epoxide hydrolase from potato (347) and purification as well as immobilization of epoxide hydrolase from rat liver (348) have been reported. Commercial availability of such enzyme preparations is a prerequisite for their application in biotransformation of fats for the preparation of products via hydration of epoxides, e.g., hydroxylated fatty acids (349).

IV. USE OF ENZYMES IN TECHNOLOGY OF OILSEEDS, OILS, AND FATS

Enzymes are gaining importance as processing aids in the technology of oilseeds, oils, and fats. Use of enzymes to facilitate the recovery of oils from oilseeds and
other oil bearing materials has become known lately. Very recently, an enzymatic process was developed for degumming of oils and fats in large-scale commercial operation.

A. Enzymes for Pretreatment of Oilseeds Prior to Oil Extraction

Conventional techniques for the recovery of oils from seeds and fruits involve grinding and conditioning by heat and moisture to disintegrate the oil-bearing cells, followed by mechanical pressing in hydraulic presses or expellers or extraction by organic solvents, such as hexane (see Chapter 8). Fats from animal tissues are frequently recovered by rendering, i.e., heat treatment with live steam. Lately, aqueous extraction processes have become known in which the seeds or fruits are ground

Figure 46 Enzymatic production of hydroxy acids by \( \omega \)-hydroxylation.

Figure 47 Enzymatic production of epoxy acids.
with water to disrupt the oil-bearing cells, followed by centrifugation to separate the oil from the solids and the aqueous phase (350).

Extensive mechanical rupturing of the cells of oil-bearing seeds is the prerequisite for efficient oil extraction (351). Several reports have suggested the use of enzymes for the rupture of the plant cell walls to release the oil contained in the cell prior to recovery of the oil by mechanical pressing, solvent extraction, or aqueous extraction. Plant cell walls are generally composed of unlignified cellulose fibers to which strands of hemicellulose are attached; the cellulose fibers are often embedded in a matrix of pectic substances linked to structural protein (352). Since substantial differences are observed in the polysaccharide composition of the cell walls of different plant species (Table 10), different combinations of cell wall–degrading enzymes (carbohydrases and proteases) have to be used for individual seeds or fruits (107,352,353). Enzymes used in cocktails for cell wall degradation include amylase, cellulase, polygalacturonase, pectinase, hemicellulase, galactomannase, and proteases (107,353).

Enzyme pretreatment followed by mechanical expelling for improved oil recovery has been used for rapeseed (107,354) and soybean (355). Typically, treatment of flaked rapeseed with commercial enzyme preparations (SP 249, Novo Nordisk

---

**Figure 48** Enzymatic production of long chain aldehydes and ketones.

**Figure 49** Preparation of dihydroxy fatty acids by epoxide hydrolase.
Figure 50  Hydration of linoleic acid monoepoxides to vic-diols by epoxide hydrolase.

Biochem North America, Inc., Franklinton, N.C., USA, and Olease, Biocon (US) Inc., Lexington, KY, USA) at 30% moisture and 50°C for 6 hours followed by drying and expelling gave 90–93% recovery of oil as compared with 72% recovery of the controls not treated with the enzymes (354).

Enzyme-assisted expeller process is now a commercial process for partial oil recovery from rapeseed (356). In the commercial process the oil release is substantially enhanced and the resulting rapeseed cake with a superior nutritional value is preferred in a number of animal feeds.

Enzyme-assisted pressing for the production of virgin grade olive oil has also been reported (107,353).

Enzyme pretreatment followed by solvent extraction for enhanced oil recovery has been used for melon seeds (357) and rapeseed (358). Incubation of autoclaved and moistened rapeseed flakes (30% moisture) with carbohydrases for 12 hours, followed by drying to 4% moisture and extraction with hexane, resulted in 4.0–4.7% enhancement of oil extraction with the different enzyme preparations in the following order: mixed activity enzyme > β-glucanase > pectinase > hemicellulase > cellulase (358). A disadvantage of the process is a rather long incubation time.

Since the earlier publication of Lanzani et al. (359) on rapeseed, sesame seed, sunflower seed, soybean, and peanut, several reports have appeared on enzyme pretreatment followed by aqueous extraction for enhanced oil recovery from coconut (352,360–363), corn germ (352,364), avocado (365,366), olives (352,367,368), mustard seed and rice bran (369), Jatropha curca seeds (370), and cocoa beans (371).

Enzyme-assisted aqueous extraction has been extensively studied on rapeseed using the mixed enzyme preparation SP-311 (Novo) and found to yield an oil with low

Table 10  Approximate Composition (%) of Cell Wall Polysaccharides of Some Oil-Bearing Materials

<table>
<thead>
<tr>
<th>Polysaccharide</th>
<th>Rapeseed</th>
<th>Coconut</th>
<th>Corn germ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pectic substances</td>
<td>39</td>
<td>—</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Mannans</td>
<td>—</td>
<td>61</td>
<td>—</td>
</tr>
<tr>
<td>Galactomannans</td>
<td>—</td>
<td>26</td>
<td>—</td>
</tr>
<tr>
<td>Arabinogalactans</td>
<td>8</td>
<td>some</td>
<td>—</td>
</tr>
<tr>
<td>Celluloses</td>
<td>22</td>
<td>13</td>
<td>39</td>
</tr>
<tr>
<td>Hemicelluloses</td>
<td>29</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Xyloglucans</td>
<td>2</td>
<td>—</td>
<td>40</td>
</tr>
<tr>
<td>Arabinobioxylans</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Others</td>
<td>2</td>
<td>—</td>
<td>10</td>
</tr>
</tbody>
</table>

Source  Ref. 352.
phospholipid content (phosphorus < 3 ppm) that can be subjected to physical refining without complex degumming procedures (352,356). The protein recovered by aqueous enzymatic extraction has higher nutritive/market value than the rapeseed obtained by conventional pressing followed by solvent extraction; however, the oil yield in aqueous enzymatic processing is somewhat lower (90–92%) compared with the conventional process (about 97.5%) (356).

Addition of plant proteases, such as papain or bromelain, to macerated fish followed by incubation at 65°C for 30 minutes and at 50–55°C for 1–2 hours, respectively, has been shown to improve the recovery of fish oils (372). Thermostable microbial proteases operating at higher temperatures (70–75°C) appear to be more suitable for this purpose because they reduce the risk of microbial growth and contamination (107).

B. Enzymatic Degumming of Oils

Degumming is an important processing step during the refining of vegetable oils, such as soybean and rapeseed (canola). Untreated vegetable oils contain varying proportions of phospholipids (phosphatides), such as phosphatidylethanolamines (PE), -inositols (PI), -ethanolamines (PE), and phosphatidic acids (PA), which impair the quality and stability of oils if they are not removed. Especially for vegetable oils that are subjected to physical refining, it is very important to eliminate the phospholipids prior to treatment of the oil with steam under vacuum for the removal of free (unesterified) fatty acids.

In the current industrial practice of degumming (373), the raw oils are treated with water or aqueous solutions of phosphoric acid, citric acid, or an alkali in order to hydrate the phospholipids that subsequently flocculate and thus can be removed by centrifugation (see also Chapter 8). Among the different classes of phospholipids PC and PI are readily hydratable, whereas both PE and PA are hydratable when combined with potassium and unhydratable when complexed with divalent metals; moreover, PA occurs as a partially dissociated acid and is nonhydratable (373,374). Phospholipase D plays an important role in the formation of nonhydratable PA. Especially during the processing of soybeans phospholipase D can remain fairly active even at a temperature of 65°C in the presence of water-saturated hexane, thereby catalyzing the hydrolysis of hydratable phospholipids, e.g., PC to nonhydratable PA; however, in aqueous media this enzyme is readily deactivated by heat (375).

Recently, Lurgi AG (Frankfurt, Germany) developed an enzymatic process (EnzyMax) for degumming of oils that involves treatment of the raw oil with phospholipase A₂ that cleaves the fatty acids esterified at the sn-2 position of the diacylglycerophospholipids yielding sn-1-acyllyso-glycerophospholipids, including sn-1-acyllyso-PE and sn-1-acyllyso-PA (Fig. 51), that can be readily hydrated and removed (374,376).

Figure 52 shows the flow sheet of the commercial EnzyMax process for degumming of oils. The enzyme used is a commercially available phospholipase A₂ (lecitase activity 10,000 IU/mL) isolated from porcine pancreas. Recently, a microbial phospholipase A₂ that is more economical than the porcine enzyme has been used successfully for degumming (376). The process consists of three stages. In the first stage, solutions of citric acid (50% w/v) and sodium hydroxide (3% w/v) are dispersed with the crude oil or water-degummed oil and the pH adjusted to approx-
Principle of enzymatic degumming of oils and fats using phospholipase A$_2$ for converting diacylglycerophospholipids to easily hydratable lysophospholipids.

\[
\begin{align*}
\text{Diacylglycerophospholipid} & \quad \xrightarrow{\text{Phospholipase A}_2} \quad \text{1-Acyl-lysoglycerophospholipid (hydratable)} \\
X & \quad \xrightarrow{\text{Choline, ethanolamine etc.}}
\end{align*}
\]

Figure 51  Principle of enzymatic degumming of oils and fats using phospholipase A$_2$ for converting diacylglycerophospholipids to easily hydratable lysophospholipids.

Approximately 5.0. The resulting mixture, maintained at a temperature of 60°C, is intimately mixed with the enzyme solution (enzyme concentrate diluted with water to 0.2% v/v) and passed through two or more enzyme reactors. The residence time of the mixture in the reactors varies from 1 to 6 hours, depending on the phosphatide content of the starting material and the quality requirement on the degummed oil. The effluent from the enzyme reactors is finally admitted to a separator to obtain the streams of degummed oil and aqueous phase containing the phosphatide sludge containing the still active enzyme that is generally reused several times by recycling into the process.

Operational data from commercial plants for degumming of 540 tons/day rape-seed oil and 400 tons/day soybean oil show a likely average consumption of 10 g enzyme concentrate per ton of oil and residual phosphorus content of the degummed oil less than 10 ppm. The enzymatically degummed oil can be subjected to physical refining after bleaching. Due to low phosphorus content and absence of soaps enzymatically degummed oils, as compared with those obtained by other degumming processes, can be bleached with standard bleaching earths instead of high activity and acidic bleaching earths. Additional economic advantages of enzymatic degum-
ming as compared with chemical neutralization and soapstock splitting are due to lower oil loss, less consumption of water, and very little effluent wastes of environmental concern.

V. PERSPECTIVES

Lipid biotechnology is a very young discipline. Although breathtaking progress has been made in genetic engineering for alteration of composition of oilseeds and oils (see Chapter 30), the commercial applications of biotechnology for the production and modification of fats and oils using whole microbial cells and isolated enzymes are restricted so far to a few areas, such as enzymatic production of cocoa butter substitutes, human milk fat replacers, and bio-ester as well as enzymatic degumming of fats and oils. Factors limiting the application of biotechnology in the area of lipids include the productivity and the cost of downstream processing in microbial processes and the price as well as the reuse properties of enzymes, such as lipases. In this regard, optimism is raised in the current literature, which contains numerous research activities aimed at development of novel lipid-based specialty products as well as mass fat products, such as feedstocks for margarines using biotechnology. With increasing production and use of enzymes in detergents one might foresee the commercial bulk scale availability of such biocatalysts at competitive prices that should facilitate their applications in wide areas of lipid biotechnology.

REFERENCES


365. M. Buenrosto and C. A. Lopez-Menguia. Enzymatic extraction of avocado oil. *Bio-


I. INTRODUCTION

Lipases (triacylglycerol acylhydrolases, EC 3.1.1.3) are enzymes that catalyze the reversible hydrolysis of triacylglycerols under natural conditions. Widely distributed in animals, plants, and microbes, lipases differ from other esterases and are unique in that their activity is greatest against water-insoluble substrates and is enhanced at the substrate (oil)–water interface; that is, they exhibit “interfacial activation.” Optimum activities are obtained in systems such as emulsions, where high surface areas of the substrate can be obtained. Lipases are active not only in normal phase emulsions where the substrate is emulsified into an aqueous system (oil-in-water), but they are also active, often more active, in invert (water-in-oil) emulsions and in reverse micelle systems containing an organic solvent solution of the substrate. Furthermore, lipases are exceedingly versatile in that they can also catalyze transesterification reactions and the stereospecific synthesis of esters, and they can act on a broad range of substrates.

Major advances have been made in our understanding of lipolytic enzymes over the past few years through solving the crystal structures of lipases from several sources. Knowledge of their structures has given insight into the mechanism of action, interfacial activation, specificity, and the nature of the active site. The high current interest in lipases, shown by the large volume of recent scientific literature on the topic, is driven by the great potential in a diversity of commercial applications for these enzymes. This chapter gives an overview of lipases and their properties, with a focus on the topics of current research interest such as lipase structure and its implications in interfacial activation and selectivity. The emphasis is on microbial lipases, particularly from fungi, with references to human pancreatic lipase for comparison.
II. LIPASES FROM DIFFERENT SOURCES

A. Animals

Lipases from various organs and tissues of several mammalian species have been investigated, but human and other pancreatic lipases are the most thoroughly studied. Pancreatic lipases are secreted into the duodenum and active on dietary triacylglycerols. They are a class of structurally similar 50-kDa glycoproteins that are characterized by their specificity toward triacylglycerols with little or no activity toward phospholipids, activation at the oil (substrate)—water interface and by colipase, and inhibition at micellar concentrations of bile salts. Pancreatic lipase can catalyze the complete breakdown of triacylglycerols to free fatty acids and glycerol. Nonhuman pancreatic lipases may differ from corresponding human lipases; for example, guinea pig pancreatic lipase differs in that it exhibits phospholipase A activity (1).

Other mammalian lipases have been studied and, while similar, they exhibit some characteristics that differ from those of pancreatic lipase. In addition to pancreatic lipase, fat digestion is aided by a series of lingual, pharyngeal, and gastric lipases that may be responsible for up to 50% of dietary fat breakdown (2). Highly stable at low pH, these enzymes are activated by bile salts and show a preference for the sn-3 position of the substrate. Human lipoprotein lipase, which functions in hydrolyzing triacylglycerols in chylomicrons and very low density lipoproteins, shows many similarities to pancreatic lipase (e.g., high sequence homology, presence of the lid). However, this lipase is not responsive to colipase but instead requires apolipoprotein C-II (apoC-II) for activity, functions as a dimer, and is activated by heparin (3–5). Hepatic lipase is confined to the liver, where it also is involved in the metabolism of lipoproteins, but it is not activated by apoC-II (6). Human milk lipase, which functions in the digestion of milk fat ingested by infants, is activated by bile salt (7).

B. Plants

Plant lipases have not received the same attention as those from other sources, but they have been reviewed recently by Mukherjee and Hills (8). Oilseed lipases have been of greatest interest among the plant lipases, and those from a variety of plant species show differences in their substrate specificity, pH optima, reactivity toward sulfhydryl reagents, hydrophobicity, and subcellular location (9). These lipases are relatively specific for the native triacylglycerols of the species from which they were isolated. They are absent from the ungerminated seed and formed during germination.

C. Microbes

There is substantial current interest in developing microbial lipases for use in biomedical and industrial applications because of their versatility and availability; in addition, they can be produced less expensively than corresponding mammalian enzymes. Hou and Johnston (10) screened 1229 bacteria, yeasts, actinomycetes, and fungi and found that about 25% were lipase positive. Microbial lipases that have received the greatest attention are inducible, extracellular enzymes having properties that are generally similar to those of human pancreatic lipase, despite differences in detail. At least some of the microbes produce a mixture of extracellular lipases formed from multiple genes (11–15) and some lipases vary by degrees of glycosyl-
Fungal lipases typically exist as monomers with molecular masses ranging from about 30 to 60 kDa. They vary in specificity, specific activity, temperature stability, and other properties; however, dimeric lipases have been reported (Table 1) (16–17).

III. PRODUCTION, ISOLATION, AND PURIFICATION

A. Lipase Production

Microbial extracellular lipase production for laboratory study is typically carried out under liquid shake culture conditions or in small fermentors. Conditions for optimum lipase production seem to be variable depending on the species. Typically, triacylglycerols (olive or soybean oil are commonly used) are placed in the culture medium to induce lipase production (17,28,29), but fatty acids may also induce lipase production (30–32). However, lipase production has been studied in the absence of lipids (e.g., with sugars as the carbon source) (33). Chang et al. (34) reported that Tween 80 and Tween 20 in the culture medium promoted lipase production and a change in the multiple forms of lipase produced by *Candida rugosa*. The following is a specific example of conditions for the maximal production of lipases by *Geotrichum candidum*: growth for 24 hours in liquid medium containing 1% soybean oil, 5% peptone, 0.1% NaNO₃, and 0.1% MgSO₄ at pH 7.0, 30°C and shaking at 300 rpm (11).

B. Isolation and Purification

Extracellular lipases have been isolated, purified, and characterized from numerous microbial species. Some examples of recently characterized lipases are illustrated in Table 1. The specific isolation and purification methodology differs in detail from study to study, but it generally involves the ammonium sulfate precipitation of proteins from the culture medium after removal of cells or mycelium, and then fractionating the proteins through a series of ion exchange, affinity, and gel filtration columns. Several recent publications can be consulted for specific procedures for isolating lipases (16–21,35,36). The following is given as an example of a lipase from *Neurospora* sp. TT-241 that was isolated and purified 371-fold (22).

1. **Ammonium sulfate fractionation.** Add solid ammonium sulfate to the clarified aqueous extract (or culture medium) up to 60% saturation. The resulting precipitate is collected by centrifugation at 12,000 × g for 30 minutes and dissolved in a minimal volume of buffer A (50 mM phosphate buffer, pH 7.0). The enzyme solution is dialyzed overnight against a 50-fold excess of the same buffer.

2. **Sephadex G-100 gel filtration chromatography.** The enzyme solution (70 mL) is applied to a Sephadex G-100 column (4 cm × 120 cm), eluted with buffer A at a flow rate of 40 mL/h, and 8 mL fractions are collected.

3. **Toyopearl phenyl-650M column chromatography.** Add ammonium sulfate to the pooled active fractions from gel filtration to a final concentration of 1 M. The enzyme solution is then applied to a Toyopearl phenyl-650M column (2.0 cm × 18 cm) that has been preequilibrated with buffer A containing 1 M ammonium sulfate. The column is washed with 300 mL.
<table>
<thead>
<tr>
<th>Organism</th>
<th>Molecular weight (kDa)</th>
<th>pH Optima</th>
<th>Optimum temperature (°C)</th>
<th>Specific activity (U/mg)</th>
<th>Specificity</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neurospora sp. TT-241</td>
<td>55</td>
<td>6.5</td>
<td>45</td>
<td>8203</td>
<td>No specificity but preferred 1- and 3-positions</td>
<td>22</td>
</tr>
<tr>
<td>Botrytis cinerea</td>
<td>60</td>
<td>6.0</td>
<td>38</td>
<td>2574</td>
<td>Highest activity with oleic acid esters</td>
<td>20</td>
</tr>
<tr>
<td>Pseudomonas fluorescens</td>
<td>33</td>
<td>8–10</td>
<td>55</td>
<td>6200</td>
<td>Nonspecific for fatty acids</td>
<td>19</td>
</tr>
<tr>
<td>Neurospora crassa</td>
<td>54</td>
<td>7.0</td>
<td>30</td>
<td>44</td>
<td>Preferred triglycerides with C_{16} and C_{18} acyl chains</td>
<td>27</td>
</tr>
<tr>
<td>Rhizopus niveus 104759 Lipase I</td>
<td>34</td>
<td>6.0–6.5</td>
<td>35</td>
<td>4966</td>
<td>—</td>
<td>23</td>
</tr>
<tr>
<td>Rhizopus niveus 104759 Lipase II</td>
<td>30</td>
<td>6.0</td>
<td>40</td>
<td>6198</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Candida parapsilosis</td>
<td>160</td>
<td>6.5</td>
<td>45</td>
<td>—</td>
<td>High specificity for long chain fatty acids, particularly polyunsaturated fatty acids</td>
<td>25</td>
</tr>
<tr>
<td>Propionibacterium acidi-</td>
<td>6–8</td>
<td>7.0</td>
<td>30</td>
<td>—</td>
<td>Preferred substrates with high saturated fatty acids</td>
<td>26</td>
</tr>
<tr>
<td>propionici</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neurospora crassa</td>
<td>54</td>
<td>7.0</td>
<td>30</td>
<td>44</td>
<td>Preferred triglycerides with C_{16} and C_{18} acyl chains</td>
<td>27</td>
</tr>
<tr>
<td>Pythium ultimum #144</td>
<td>270</td>
<td>8.0</td>
<td>30</td>
<td>63</td>
<td>1,3-Specific, preferring substrates with higher unsaturation</td>
<td>17</td>
</tr>
<tr>
<td>Rhizopus delemar ATCC 34612</td>
<td>30.3</td>
<td>8.0–8.5</td>
<td>30</td>
<td>7638</td>
<td>—</td>
<td>18</td>
</tr>
<tr>
<td>Fusarium heterosporium</td>
<td>31</td>
<td>5.5–6.0</td>
<td>45–50</td>
<td>2010</td>
<td>1,3-Specific; preferred triacylglycerols with C_{6}–C_{12} fatty acids</td>
<td>21</td>
</tr>
<tr>
<td>Penicillium roquefortii</td>
<td>25</td>
<td>6.0–7.0</td>
<td>30</td>
<td>4063</td>
<td>Preferred triacylglycerols with C_{17}–C_{18} fatty acids</td>
<td>16</td>
</tr>
<tr>
<td>Penicillium sp. uzim-4</td>
<td>27</td>
<td>7.0</td>
<td>25</td>
<td>1001</td>
<td>1,3-Specific, discriminates against diglycerides and active at low surface pressures</td>
<td>24</td>
</tr>
</tbody>
</table>
of buffer A containing 1 M ammonium sulfate and then eluted with 1 L of a linear gradient from 1 to 0 M ammonium sulfate in buffer A at a flow rate of 60 mL/h; 4 mL fractions are collected.

4. *Ultrogel-HA hydroxyapatite column chromatography.* The pooled active fractions from the Toyopearl column are further purified by passage through an Ultrogel-HA column (3.0 cm × 9 cm), preequilibrated with 10 mM phosphate buffer (pH 7.0). After being washed with 10 mM phosphate buffer, the column is eluted with a 1 L linear gradient from 10 to 500 mM phosphate buffer at a flow rate of 50 mL/h, and 5 mL fractions are collected. The active fractions are pooled and stored at −20°C.

These procedures are carried out at 4°C. The purity of lipase active fractions is determined by sodium dodecyl sulfate polyacrylamide gel electrophoresis.

**IV. ASSAY OF LIPASES**

**A. Review of Emulsion Systems**

Lipases are assayed in emulsions where high substrate surface areas can be achieved. Since, in addition to high interfacial area, the chemical and physical environments at the substrate–water interface are important to the adsorption and activity of lipases, it is useful to understand the nature of the different types of emulsion used for lipase assay or in various applications. Emulsions are mixtures of two immiscible liquids (e.g., oil and water); one of the components is dispersed as very small droplets, or particles, and the mixture is stabilized by a surface-active agent, or surfactant. Emulsions are classified as macro- or microemulsions where the dispersed particles are either greater or smaller than one micrometer in diameter, respectively. Macroemulsions are turbid, milky in color, and thermodynamically unstable (i.e., they will ultimately separate into the two liquid phases). On the other hand, microemulsions are homogeneous and stable.

Depending on the conditions of formation, and particularly the nature of the surfactant, macroemulsions may be normal phase (oil-in-water) or reversed phase (water-in-oil), invert. In the former, the oil is emulsified into the aqueous phase, and the surfactant forms a monolayer film around the dispersed oil droplets, whereby the hydrophobic moiety of the surfactant extends into the oil and the polar moiety is at the droplet surface (Fig. 1). In reversed phase emulsions, the aqueous phase is dispersed in the oil with the orientation of the surfactant molecules reversed (Fig. 1).

Amphiphilic (surfactant) molecules undergo self-organization into spheroidal particles when dissolved in certain organic solvents such as isooctane with the polar head groups oriented inward and hydrophobic tails outward. These particles are referred to as reverse micelles and are less than 1 μm in diameter (Fig. 1). Water can be “solubilized” in the organic solvent by becoming entrapped in the particles at up to several dozens of molecules per molecule of surfactant. Typically, reverse micelles are formed when the molar ratio of water to surfactant is less than 15, which is expressed as water activity, \( w_0 \) (i.e., molarity of water to molarity of surfactant) or \( R \). Enzymes such as lipases can be entrapped within the aqueous phase particles of the invert emulsions or reverse micelles of microemulsions, where they retain their activity. In the latter case, the enzymes are isolated from the organic solvent. Luisi
et al. (37) and Sanchez-Ferrer and Garcia-Carmona (38) can be consulted for more information on microemulsions.

B. Normal Phase Emulsions

Most lipase assays for hydrolytic reactions are carried out in normal phase emulsions whereby the water-insoluble oil substrate is emulsified by sonication into a buffered aqueous enzyme preparation. The emulsion is stabilized with an emulsifier and the aqueous phase often contains Ca\(^{2+}\). The specific ingredients, buffer types, pH, and relative amounts of components vary widely with the author and specific lipase being assayed. An example of such an assay is as follows: 2 mL 0.2 M Tris maleate–NaOH buffer (pH 8.2), 1 mL 0.03 M Ca\(_2\)Cl\(_2\), 5 mL distilled water, 1 mL olive oil, and 1 mL enzyme solution. The emulsion is incubated at 30\(^\circ\)C for 60 minutes or an appropriate time depending on lipase activity, and the reaction is stopped with 20 mL of acetone ethanol (1:1). Common emulsifiers used in lipase assays include polyvinyl alcohol and gum arabic at 1–2% by volume of assay mixture. Hydrolytic activity is most often determined by titration of the fatty acid products of the reaction with NaOH. Other methods and assay conditions have been reviewed by Jensen (39).

C. Invert Emulsions

Using lipases from several fungi, Mozaffar and Weete (40) reported a reaction mixture containing 5 mL olive oil, 0.1 mL 520 mM taurocholic acid in 50 mM sodium phosphate buffer (pH 7.5), and 0.1 mL enzyme preparation. Final concentrations of taurocholic acid and water in the 5.2 mL of reaction mixture were 10 mM and 4%. The mixture was emulsified by vortexing for about 30 seconds and incubated at 45\(^\circ\)C.
without shaking for 30 minutes, whereupon the reaction was stopped as described above. Up to 90% of the substrate was hydrolyzed under these conditions by \textit{C. rugosa} lipase when the incubation time was extended to 48 hours with periodic additions of buffer (0.2 mL) during the incubation.

**D. Microemulsion: Reverse Micelles**

1. **Hydrolysis**

The desired amount of concentrated buffered lipase solution is poured into 5 mL of 50 mM AOT [Aerosol–OT, bis (2-ethylhexyl) sodium sulfosuccinate]–isooctane solution containing 10% v/v of the substrate. The amount of enzyme solution depends on the \textit{R} value (e.g., 10.5). The reaction is initiated by vortexing until clear, and the mixture is incubated at 30°C for 15 minutes; then 0.4 mL is added to 4.6 mL benzene and 1.0 mL cupric acetate–pyridine solution, and the reaction is stopped by vortexing (41a). Fatty acids liberated by the hydrolytic reaction are determined according to Lowry and Tinsley (42). Other examples of studies involving hydrolysis by lipases in organic solvent–reverse micelle systems are cited in Table 2 (41a, 41b, 43–78).

2. **Transesterification**

Using a lipase from \textit{Rhizopus delemar}, Osterberg et al. (79) used the following system for transesterifying triacylglycerols with stearic acid: isooctane (91.65 wt %) was mixed with AOT at 100–200 mM, aqueous 0.066 M phosphate buffer (pH 6) (1.0%), and substrate (5.0%). The enzyme in the buffer was used at 1.5 U/mg substrate. The reaction was carried out at 35°C under nitrogen with magnetic stirring, and was stopped by raising the temperature to 100°C and holding at that temperature for 10 minutes. Other examples of studies involving transesterification and synthetic reactions in organic solvent–reverse micelle systems are cited in Table 2.

**V. PROPERTIES AND REACTIONS**

**A. Structure**

Over the past several years, the crystal structures of several mammalian and microbial lipases have been determined. Generally, lipases are \textit{\alpha/\beta} proteins with a central core of a mixed \textit{\beta} sheet containing the catalytic triad composed of Ser\cdots His\cdots Asp, and a surface loop restricting access of the substrate to the active site.

1. **Animal Lipases**

Human pancreatic lipase is folded into two domains, a larger N-terminal domain comprising residues 1–335 and a smaller C-terminal domain (residues 336–449). The core of the N domain is formed by a nine-stranded, \textit{\beta}-pleated sheet in which most of the strands run parallel to one another. Seven \textit{\alpha}-helical segments of varying length occur in the strand connections, and six of them pack against the two faces of the core sheet. The C domain is formed by two layers of antiparallel sheets, the strands of which are connected by loops of varying length. The N domain contains the active site, a glycosylation site, a Ca\textsuperscript{2+}-binding site, and possibly a heparin-binding site. The active site is buried beneath a short amphipathic \textit{\alpha}-helical surface loop, termed the “flap” or “lid” (80,31). Colipase (see below), binds exclusively to
<table>
<thead>
<tr>
<th>Reaction type</th>
<th>Topic</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Synthesis</strong></td>
<td>Wax ester synthesis in a membrane reactor with lipase–surfactant complex in hexane</td>
<td>55</td>
</tr>
<tr>
<td></td>
<td>Formation of polyol–fatty acid esters</td>
<td>56</td>
</tr>
<tr>
<td></td>
<td>Esterification reactions catalyzed by lipases in microemulsions: role of enzyme localization in relation to selectivity</td>
<td>57</td>
</tr>
<tr>
<td></td>
<td>Esterification of oleic acid with glycerol in monolayer and microemulsion systems</td>
<td>58</td>
</tr>
<tr>
<td></td>
<td>Synthesis of sugar esters</td>
<td>59</td>
</tr>
<tr>
<td></td>
<td>Polyunsaturated fatty acid glyceride synthesis</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td>Monoacylation of fructose</td>
<td>61</td>
</tr>
<tr>
<td></td>
<td>Double enantioselective esterification of racemic acids and alcohols</td>
<td>62</td>
</tr>
<tr>
<td></td>
<td>Esterification of oleic acid and methanol in hexane</td>
<td>63</td>
</tr>
<tr>
<td></td>
<td>Esterification of glycerol: synthesis of regioisomerically pure 1,3-sn-diacylglycerols and monoacylglycerols</td>
<td>47,48</td>
</tr>
<tr>
<td></td>
<td>Synthesis of DHA-rich triglycerides</td>
<td>64</td>
</tr>
<tr>
<td></td>
<td>Synthesis of acylated glucose</td>
<td>53</td>
</tr>
<tr>
<td></td>
<td>Synthesis of mono- and diglycerides</td>
<td>49</td>
</tr>
<tr>
<td><strong>Interesterification</strong></td>
<td>Interesterification of butterfat</td>
<td>65</td>
</tr>
<tr>
<td></td>
<td>Interesterification of triglycerides</td>
<td>66</td>
</tr>
<tr>
<td></td>
<td>Enzymatic interesterification of triolein in canola lecithin–hexane reverse micelles</td>
<td>67</td>
</tr>
<tr>
<td></td>
<td>Alteration of melting point of tallow–rapeseed oil</td>
<td>68</td>
</tr>
<tr>
<td></td>
<td>Interesterification of phosphatidylcholine</td>
<td>69</td>
</tr>
<tr>
<td></td>
<td>Incorporation of long chain fatty acids into medium chain triglycerides</td>
<td>46</td>
</tr>
<tr>
<td></td>
<td>Kilogram-scale ester synthesis</td>
<td>70</td>
</tr>
<tr>
<td></td>
<td>Incorporation of n-3 polyunsaturated fatty acids into vegetable oils</td>
<td>71</td>
</tr>
<tr>
<td></td>
<td>Incorporation of EPA and DHA into groundnut oil</td>
<td>45</td>
</tr>
<tr>
<td></td>
<td>Interesterification of milk fat with oleic acid</td>
<td>72</td>
</tr>
<tr>
<td></td>
<td>Incorporation of exogenous DHA into bacterial phospholipids</td>
<td>73</td>
</tr>
<tr>
<td></td>
<td>Interesterification of triglycerides and fatty acids by surfactant-modified lipase</td>
<td>74</td>
</tr>
<tr>
<td></td>
<td>Transesterification of rapeseed oil and 2-ethylhexanol</td>
<td>75</td>
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<tr>
<td></td>
<td>Transesterification of cocoa butter</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td>Modification of phospholipids</td>
<td>54</td>
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<tr>
<td></td>
<td>Diacylglycerol formation</td>
<td>50</td>
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<tr>
<td></td>
<td>Transesterification of palm oil</td>
<td>76</td>
</tr>
<tr>
<td><strong>Hydrolysis</strong></td>
<td>Hydrolysis of phosphatidylcholine by an immobilized lipase</td>
<td>43</td>
</tr>
<tr>
<td></td>
<td>Palm kernel olein hydrolysis</td>
<td>77</td>
</tr>
<tr>
<td></td>
<td>Olive oil hydrolysis</td>
<td>41a,41b</td>
</tr>
<tr>
<td></td>
<td>Production of polyunsaturated fatty acid enriched fish oil</td>
<td>78</td>
</tr>
<tr>
<td></td>
<td>Enrichment of γ-linolenic acid from evening primrose oil and borage</td>
<td>47,48</td>
</tr>
</tbody>
</table>

*The references cited are representative; they are not intended to be inclusive.*
The C domain of the protein through hydrophobic interactions and ion pairing (82), and the binding does not induce a conformational change (83–85). Calcium ions activate and reduce the lag phase of human pancreatic lipase, particularly for mixed bile acid–lipase complexes (86).

The structures of pancreatic lipase from other systems have been recently reported, including guinea pig (1) and horse (87). Overall, the nonhuman pancreatic lipases are structurally similar to that from humans but may differ in detail. For example, the guinea pig pancreatic lipase does not possess the lid that is typical of other lipases (1). Also, unlike the pancreatic lipases from humans and pigs, the horse enzyme is not glycosylated (87).

2. Fungal Lipases

The crystal structures of several fungal lipases have also been determined: for example, Rhizomucor miehei (88,89), G. candidum (90,91), C. rugosa (92), C. antarctica (93), and Humicola lanuginosa (94). Although there are no obvious sequence similarities between pancreatic and fungal lipases, except for the Gly-X-Ser-X-Gly consensus sequence in the active site region (87,95), there are structural similarities. Fungal lipases are α/β proteins and have similar topologies based on a large central mixed β-pleated sheet pattern, mainly parallel, but the connectivities between strands vary. The two lipases (see below) of G. candidum have 554 amino acids and share about 85% sequence homology (13,96). Also, many fungal lipases possess the serine protease catalytic triad Ser···His···Asp in their active sites; however, glutamic acid is substituted for aspartic acid in the lipases from G. candidum (90) and C. rugosa (92). Lipase II gene from G. candidum has been cloned (13,97a), and the results of probing the active site by site-directed mutagenesis are consistent with X-ray crystallography data in that the Ser···His···Glu is the active site (97a).

Fungal lipases also possess a lid that prevents access of the substrate to the active site, form a functional oxyanion hole, and have an interfacial binding site (Gly-X-Ser-X-Gly), but the motion of the lid differs between fungal and pancreatic lipases. Lid rearrangements by human pancreatic lipase (84,85), C. rugosa (92), and G. candidum (90) involve more than one loop. The lid is closed and covers the active site in G. candidum and is open in C. rugosa; otherwise lipases from these two sources are very similar in sequence and structure. The three-dimensional structure of the extracellular lipase from Rhizopus delemar based on X-ray crystallographic coordinates is shown in Figure 2, with the lid partially open exposing the active site residues Ser145, Asp204, and His257 (Fig. 2A) and residues 86–92 in the partially closed conformation (Fig. 2B) (97b).

3. Bacterial Lipases

Lipases from several Pseudomonas species have been crystallized, and preliminary X-ray crystallographic analyses of such species as P. cepacia (98), P. glumae (99), and P. fluorescens (100a) have been conducted. Their molecular masses, which are in the 30- to 35-kDa range, show extensive sequence homology to one another but little to those of other lipases; the common G-X-S-X-G sequence in the active site region is conserved, and activity appears at the substrate–water interface. On the other hand, lipases from Bacillus species have molecular weights at about 19 kDa, have the sequence A-X-S-X-G instead of the characteristic sequence, and do not exhibit interfacial activation (100b).
B. Interfacial Activation and the Hydrolytic Reaction

In 1958 Sarda and Desnuelle (101) showed that pancreatic lipase does not exhibit normal Michaelis–Menten kinetics with respect to substrate concentration. Lipases are inactive in aqueous media with the substrate present in its monomeric form, but there is a sharp increase in activity when the substrate exceeds the critical micelle concentration. The inactive enzyme must first adsorb to the surface of the bulk substrate, which initiates interfacial activation (Fig. 3). Interfacial enzyme kinetics in lipolysis have been reviewed by Verger and Haas (102a).

Solving the crystal structures of lipases has given insight into the mechanism of interfacial activation (102b). It is believed that the preferred conformation in aqueous solution is with the lid covering the active site, thus denying access to the substrate. Adsorption of the lipase to the interface involves a conformational change in the enzyme whereby the lid, which covers a cavity containing the active site and is held in place by mostly hydrophobic and some hydrogen bonds, undergoes re-orientation. This is accompanied by additional conformational changes that expose...
Figure 2  (Continued)

Figure 3  Schematic representation of the adsorption and hydrolytic activities of lipases at the oil (below the plane) and aqueous (above the plane) interface: E, enzyme in the aqueous phase; Ea, enzyme adsorbed at the interface; Ea*, activated enzyme; S, Substrate; Ea*S, enzyme–substrate complex; Ea*–Ac, acylenzyme; P1, product (diacylglycerol); P2, product (fatty acid).
the active site and a larger hydrophobic site, and allow access of the substrate to the active site. The working hypothesis for the mechanism of interfacial activation is based on the three-dimensional structural analyses of lipases from *Rhizomucor miehei* (88,103), human pancreatic lipase (81), *G. candidum* (90,91), and *C. rugosa* (92,104). The two latter lipases, for example, are globular, single-domain proteins built around an 11-stranded mixed $\beta$ sheet with domains of $45 \times 60 \times 65$ Å (105).

Activation of *R. miehei* lipase involves the movement of a 15 amino acid long lid in a hinge-type, rigid-body motion that transports some of the atoms of a short $\alpha$ helix by more than 12 Å (103). This, combined with another hinge movement, results in the exposure of a hydrophobic area representing 8% of the total molecular surface. In *C. rugosa*, comparison of open (92) and closed (104) conformation indicates that activation of the lipase requires the movement and refolding, including a cis-to-trans isomerization of a proline residue, of a single surface loop to expose a large hydrophobic surface where the substrate likely interacts. Lid reorganization contributes to the formation of a catalytically competent oxyanion hole and creation of a fully functional active site (105). The scissile fatty acyl chain is bound in a narrow, hydrophobic tunnel, where modeling studies suggest that the substrate must adopt a tuning fork conformation (105,106). There is a tryptophan residue at the tip of the lid (Trp$^{89}$) in the lipase from *H. lanuginosa* that plays an important role in hydrolytic activity (107). When Trp$^{89}$ is substituted with other amino acids, activity drops substantially and variably depending on the substituent amino acid.

It is difficult to visualize how the hydrophobic substrate molecule, which is buried in the oil surface, gains access to the active site, which is buried within the water-soluble lipase molecule. Blow (108) has suggested that upon activation, a "hydrophobic seal" forms at the interface that allows the substrate to enter the active site without interacting with the bulk water; that is, the enzyme partially withdraws the substrate molecule from the bulk oil with at least some of the acyl chains projecting into the lipid.

Cygler et al. (105) have postulated that there are two tetrahedral intermediates in the lipase-catalyzed hydrolysis of esters. Formation of a noncovalent Michaelis complex between the lipase and triacylglycerol is followed by formation of a tetrahedral, hemiacetal intermediate resulting from a nucleophilic attack by the serine O$^\ominus$. The oxyanion resulting from formation of the enzyme–substrate complex is stabilized by the amide groups of the oxyanion hole (e.g., Gly$^{123}$ in *C. rugosa*) and $\alpha$ helix following the active site serine (105). Intermediate formation is followed by cleavage of the substrate ester bond, breakdown of the tetrahedral intermediate to the acyl enzyme, and protonation and dissociation of the diacylglycerol. The serine ester of the acylated enzyme is attacked by an activated water molecule to form a second tetrahedral intermediate, which is cleaved to give rise to the protonated enzyme (serine residue) and fatty acid.

### C. Activation/Inhibition

Calcium may stimulate lipase-catalyzed hydrolytic activity by (a) binding to the enzyme resulting in a change in conformation, (b) facilitating adsorption of the lipase to the substrate–water interface, and/or (c) removing from the interface fatty acid products of hydrolysis that may reduce end-product inhibition of the reaction. Activation of human pancreatic lipase by calcium is complex and variable depending
on the substrate and presence or absence of bile acids (86). Calcium effects on microbial lipase activity may be variable depending on the enzyme source and assay conditions. For example, stimulation of *C. rugosa* lipase activity was attributed to the formation of calcium salts of fatty acid products in a normal phase emulsion, with olive oil as the substrate but not with tributyrin; however, calcium had no effect in an invert emulsion (109). In a nonemulsion system (i.e., without emulsifier), calcium had no effect on *C. rugosa* lipase activity with olive oil as the substrate but tended to offset the inhibitory effects of bile acid (Mozaffar and Weete, unpublished).

Human pancreatic lipase is inhibited by the bile salts, and the inhibition can be overcome by the 10-kDa protein colipase. Bile salt coating of the substrate micelles creates a negatively charged surface that is believed to inhibit adsorption of the bile salt–lipase complex to the interface (110). Colipase overcomes the inhibitory effect of the bile salts through formation of a 1:1 complex with lipase that facilitates adsorption at bile-salt-coated interfaces (83,110). Naka and Nakamura (111) found that although the bile salt sodium taurodeoxycholate inhibited pancreatic lipase activity when tributyrin was the substrate, a result that has been widely reported and cited by others, and colipase could reverse the inhibition, the bile salt actually stimulated hydrolytic activity when triolein was the substrate. This was attributed to the fact that triolein is a more natural substrate for the lipase than tributyrin. On the other hand, when sodium taurocholate was added to an emulsion assay of the lipase from *C. rugosa* with olive oil as the substrate, activity was progressively inhibited from 0.1 mM to 0.8 mM concentration of the bile salt (112). Relatively high activity at the lowest concentration was attributed to the role of the bile salt in the stabilization of oil particles in the emulsion and providing high interfacial area for adsorption of the lipase, and inhibition was due to interaction with the enzyme such that adsorption was reduced.

A variety of substances have been shown to inhibit lipase activity; examples include anionic surfactants, certain proteins, metal ions, boronic acids, phosphorus-containing compounds such as diethyl p-nitrophenyl phosphate, phenylmethyl sulfonylfluoride, certain carbamates, β-lactones, and diisopropylfluorophosphate (113).

### D. Selectivity

Lipases can be separated into three groups according to specificity (114,115). The first group shows no marked specificity with respect to the position of the acyl group on the glycerol molecule, or to the specific nature of the fatty acid component of the substrate. Complete breakdown of the substrate to glycerol and fatty acids occurs with nonspecific lipases. Examples of such lipases are those from *C. (cylindraceae) rugosa, Corynebacterium acnes*, and *Staphylococcus aureus*. The second group attacks the ester bonds specifically at the 1- and 3-positions of the substrate, with mixtures of di- and monoacylglycerols as products. Because of the instability of intermediate 1,2-di-, 2,3-di-, and 2-monoacylglycerols (i.e., migration of the fatty acid from the 2-position to the 1- or 3-position), these lipases may catalyze the complete breakdown of the substrates. Most microbial lipases fall into this group; examples include those from *Aspergillus niger, Rhizopus delemar (oryzae), R. miehei*, and *Mucor javanicus*. Members of the third group of lipases show preference for a specific fatty acid or chain length range, and are less common. The most widely studied lipase in this regard is that from *G. candidum*, which shows specificity for
long chain fatty acids with a cis double bond in the C-9 position (see below) (116). Other lipases that show some preference for specific fatty acids are those from *C. rugosa* (C18:1 *cis*-9), strains of *A. niger* (C10 and C12 or C18:1 *cis*-9), *M. miehei* (C12), *Rhizopus arrhizus* (C8, C10) (115), *Humicola lanuginosa* #3 (117), and human gastric lipase (C8 and C10) (118). A lipase from *Penicillium camembertii* hydrolyzes only mono- and diacylglycerols (119).

Negative selectivity has been observed where the lipase from *C. cylindracea* (*rugosa*) selected against triacylglycerol molecules containing DHA (120); *G. candidum* lipase preparations have shown similar discrimination for γ-linolenate in borage oil (121), and erucic acid from rapeseed oil (122).

Lipase selectivity/specificity may be due to structural features of the substrate (e.g., fatty acid chain length, unsaturation, stereochemistry), physicochemical factors at the interface, and/or differences in the binding sites of the enzyme. Stereoselectivity of enzymes can be influenced by temperature and hydrophobicity of the solvent. Recently, Rogalska et al. (123) showed that the enantioselectivity of lipases from *R. miehei*, *C. antarctica* B, lipoprotein lipase, and human gastric lipase toward monolayers of racemic dicaprin was enhanced at low surface pressures, while catalytic activity decreased.

Variations in specificity of lipase preparations from different fungi, different strains of the same species, and the same strain cultured under different conditions may be due to the production of multiple isoforms with differing specificities (124). The lipase(s) of *G. candidum* have been of particular interest because those from some strains exhibit a relatively high preference for ester bonds involving fatty acids with a *cis*-9 double bond (e.g., oleic acid). *Geotrichum* species and strains produce at least two glycosylated lipases, most often designated lipase I and II, or less often A or B, which are coded for by two genes (13,14,96,125). Additional isoforms may be produced that differ by degrees of glycosylation. There are conflicting reports on the specificities of the two lipases. For example, of the two lipases produced by *G. candidum* CMICC 335426, lipase B showed high specificity for the *cis*-9 unsaturates whereas lipase A of this strain and, according to Sidebottom et al. (15), lipases I and II of ATCC 34614, showed no preference. Bertolini et al. (125) recently cloned lipases I and II from *G. candidum* ATCC 34614 into *Saccharomyces cerevisiae*, then isolated and purified the two lipases from this yeast and determined their substrate specificities. Lipase I showed higher specificity than lipase II for long chain unsaturated fatty acyl chains with *cis*-9 double bond, and lipase II showed a preference for substrates having short acyl chains (C8 to C14). These investigators also showed that sequence variation in the N-terminal amino acids of these lipases, or the lid, does not contribute to variation in substrate preference. Not all strains of *G. candidum* exhibit the preference for unsaturated fatty acids (11). The specificities of isoforms of lipases from *Geotrichum* are shown in Table 3 (15,126–129).

Cygler et al. (105) suggested that the basis for the selectivity differences between lipases I and II from *G. candidum* involves key amino acid residues along the internal cavity presumed to be the binding site of the scissile acyl chain and that selectivity does not involve the lid. Studies with a cloned lipase from *R. delemar* (130) appear to support this suggestion. For example, substitution of certain amino acids (e.g., Phe → Asp) through site-directed mutagenesis in the substrate-binding region resulted in an almost twofold increase in the preference for tricaprylin relative.
Table 3  Selectivities of Multiple Extracellular Lipases from *Geotrichum* Species/Strains and *Mucor miehei*

<table>
<thead>
<tr>
<th>Fungus/strain</th>
<th>Molecular mass (kDa)</th>
<th>Selectivity</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hydrolysis</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>G. candidum</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ATCC 34614</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lipase I</td>
<td>50.1</td>
<td>No preference for C18:1</td>
<td>15</td>
</tr>
<tr>
<td>Lipase II</td>
<td>55.5</td>
<td>No preference for C18:1</td>
<td></td>
</tr>
<tr>
<td><em>G. candidum</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CMICC 335426</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lipase A</td>
<td>53.7</td>
<td>Preference for C16:0 relative to C18:1Δ2</td>
<td>9</td>
</tr>
<tr>
<td>Lipase B</td>
<td>48.9</td>
<td>Selectivity for fatty acids with cis-9 double bond</td>
<td></td>
</tr>
<tr>
<td><em>G. candidum</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ATCC 66592</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lipase I</td>
<td>61</td>
<td>Hydrolyzed C16:0 methyl ester at 60% of initial</td>
<td>126</td>
</tr>
<tr>
<td></td>
<td></td>
<td>velocity of that of C18:1 methyl ester</td>
<td></td>
</tr>
<tr>
<td>Lipase II</td>
<td>57</td>
<td>Hydrolyzed C16:0 methyl ester at initial velocity</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>that was only 7% of that of C18:1 methyl ester</td>
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</tr>
<tr>
<td><em>G. candidum</em></td>
<td></td>
<td></td>
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<tr>
<td>ATCC 34614</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lipase I</td>
<td>64</td>
<td>Showed high preference for triolein and TAG</td>
<td>127</td>
</tr>
<tr>
<td></td>
<td></td>
<td>with C-8</td>
<td></td>
</tr>
<tr>
<td>Lipase II</td>
<td>66</td>
<td>Same as lipase I</td>
<td></td>
</tr>
<tr>
<td><em>Geotrichum</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>sp. F0401B</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Lipase A</td>
<td>62</td>
<td>Incompletely 1,3-specific toward triolein but</td>
<td>128</td>
</tr>
<tr>
<td></td>
<td></td>
<td>able to hydrolyze at 2-position at slower rate</td>
<td></td>
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<tr>
<td>Lipase B</td>
<td>—</td>
<td>Nonspecific positional specificity (presumed</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>mixture of A and C)</td>
<td></td>
</tr>
<tr>
<td>Lipase C</td>
<td>58</td>
<td>Incompletely 2-specific</td>
<td></td>
</tr>
<tr>
<td><strong>Synthesis</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>G. candidum</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NRRL Y-553</td>
<td></td>
<td>Reactivities slow for γ-linolenic and ricinoleic</td>
<td>129</td>
</tr>
<tr>
<td></td>
<td></td>
<td>acids relative to oleic acid; oleic acid</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>esterifies 2.5 times faster than C16:0 to 1-</td>
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<tr>
<td></td>
<td></td>
<td>butanol, but 50 times faster with 2-methyl-1-</td>
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<tr>
<td></td>
<td></td>
<td>propanol or cyclopentanol</td>
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</tbody>
</table>

At least five genes encoding lipases have been found in *C. cylindracea (ruga-*)*, but the individual isoforms have not been purified and had their specificities determined (132). Based on gene sequences, the lipase genes of *C. cylindracea* and *G. candidum* appear to belong to the cholinesterase family.

### E. Immobilization

One of the limitations to the industrial uses of lipases is cost effectiveness. This can be improved by reuse of the lipase which can be accomplished by immobilizing the enzyme on an inert support. Immobilized lipases have been studied for hydrolytic (12,43,133–138), synthetic (139–143), and inter-/transesterification (144–147) reactions in both aqueous and organic media. Materials tested as solid supports for
lipases include ion exchange resins (DEAE-Sephadex A50 or Amberlite IRA94) (137) and others (134), adsorbants such as silica gel (143), microporous polypropylene (133), and nylon (148). In other cases, lipases have been immobilized by entrapping them in gels of photo-cross-linkable resins (ENT and ENTP) (135). Immobilization using methods described in the references cited above does not involve chemical modification of the enzyme. More recently, Braun et al. (141) achieved immobilization of C. rugosa lipase on nylon by covalent attachment to the support after conversion of carbohydrate groups on the enzyme to dialdehydes.

VI. INDUSTRIAL APPLICATIONS

A. General Uses

The current industrial enzyme market is about $600 million, with lipases representing about 4% of the worldwide market. The three major industrial enzyme companies worldwide are Novo Nordisk (>50%), Genencor (35%), and Solvay. Lipases are currently used, or have the potential for use, in a wide range of applications: in the dairy industry for cheese flavor enhancement, acceleration of cheese ripening, and lipolysis of butterfat and cream; in the oleochemical industry for hydrolysis, glycerolysis, and alcoholysis of fats and oils; and for the synthesis of structured triglycerides, surfactants, ingredients of personal care products, pharmaceuticals, agrochemicals, and polymers (149,150).

The Colgate–Emery process, currently used in the steam fat-splitting of triacylglycerols, requires 240–260°C and 700 psi, has energy costs, and results in an impure product requiring redistillation to remove impurities and degradation products. Also, this process is not suitable for highly unsaturated triacylglycerols (150). Lipase-catalyzed reactions offer several benefits over chemical reactions, including stereospecificity, milder reaction conditions (room temperature, atmospheric pressure), cleaner products, and reduced waste materials (44,151,152).

The largest current use of industrial enzymes is in laundry detergents, where they combine environmental friendliness and biodegradability with a low energy requirement and efficiency at low concentrations. The current U.S. market share of enzymatic laundry detergents is approaching 80%, and the U.S. detergent enzyme market is about $140 million. Essentially four types of enzyme are used in detergents: proteases, amylases, lipases, and cellulases. These enzymes perform multiple functions (e.g., stain removal, antiredeposition, whiteness/brightness retention, and fabric softening). Proteases were the first and are the most widely used enzymes in detergent formulations. Lipases are relatively new introductions to detergents, where they attack oily and greasy soils and contribute to making the detergents particularly effective at lower wash temperatures. However, a current limitation is that most lipases are unstable in alkaline conditions in the presence of anionic surfactants used in laundry detergents (19). However, some lipases may be relatively resistant to certain surfactants (153).

B. New Lipases/Modification of Known Lipases

Early studies with fungal lipases focused on the isolation and characterization of extracellular lipases from various species. Some of the thoroughly studied fungal lipases include those from C. (cylindracea) rugosa, R. miehei, P. camembertii, H.
lanuginosa, C. antarctica B, Rhizopus delemar, and G. candidum, all of which are commercially available. Many of these lipases have relatively high specific activities: 3485 U/mg for Mucor miehei lipase A (12) and 7638 U/mg for Rhizopus delemar (18) (see Table 1 for other examples). Lipases that have received the most attention are mainly those having relatively high activities or certain properties that make them commercially attractive. Other than additional strains of known lipase producers, there seems to be no pattern among the fungi or yeasts from a taxonomic point of view that would direct future studies on where to find prolific lipase producers or lipases with specific properties.

Lipases have been modified using either chemical or molecular approaches to alter their properties and to identify structure–activity relationships. For example, lipases have been chemically modified with polyethylene glycol to render them more soluble in organic media. Recently, Kodera et al. (154) produced amphipathic chain-shaped and copolymer derivatives of lipases from Pseudomonas fragi or P. cepacia that were soluble in aqueous and hydrophobic media and exhibited catalytic activities for esterification and transesterification reactions, as well as for hydrolysis. The modified lipase showed preference for the R isomer of secondary alcohols in esterification reactions.

Molecular approaches have been used to increase the production of a lipase from the fungus Rhizopus delemar (130). The gene for this lipase codes for a preproenzyme that is posttranslationally modified to the mature enzyme. A cloned cDNA for the precursor polypeptide of the lipase (155) was altered by site-directed mutagenesis to produce fragments that code for the proenzyme and mature enzyme (130). When inserted into E. coli BL21 (DE3), the quantities of lipase from a 1-L culture exceeded those obtained from the fungal culture by 100-fold. Other examples of gene modification of lipases are given in Sec. V.D.

C. Production Synthesis/Modification

There are many examples of uses for lipases in product synthesis/modification. One of the major areas of interest is in the use of lipase-catalyzed interesterification to improve the nutritional value, or alter the physical properties, of vegetable or fish oils. This is achieved, for example, by increasing the content of docosahexaenoic acid (DHA) or eicosapentaenoic acid (EPA) of these oils. These long chain ω3 (n-3) fatty acids have been incorporated into several vegetable oils using a lipase from Mucor miehei (45,156), medium chain triglycerides (46), and cod-liver oil (157). The n-3 fatty acid content of menhaden and anchovy oils (158) and tuna oil (120,159) has also been increased by lipase-catalyzed interesterification. Another fatty acid of interest is γ-linolenic acid (GLA), which is applicable in a wide range of clinical disorders. GLA has been enriched in evening primrose and borage oils by several fungal lipases (47,48).

Other research involving synthesis/modification includes the synthesis of mono- and diglycerides (49,50) including regioisomerically pure products (51,52), synthesis of acetylated glucose (53), modification of phospholipids into biosurfactants (54), hydrolysis of phosphatidylcholine (43), and production of high value specialty fats such as cocoa butter substitutes or hardened vegetable oils with butterfat properties (151). The production of high-value fats takes advantage of the 1,3-specificity of lipases that could not be achieved by chemical synthesis (44). Some recent examples of research involving synthesis/modification by lipases were given in Table 2.
VII. SUMMARY

Lipases are exceedingly interesting enzymes because the relationship between their structure and activity presents an intellectual challenge and because their versatility offers a broad range of possible industrial applications. However, interest in the lipases has begun to move from academic curiosity to full commercialization in terms of the availability of lipases and their industrial use. For example, 50 tons per year of the chiral intermediate methyl methoxyphenyl glycidate is produced based on a lipase-catalyzed process (141).

Although lipases have high potential for a variety of industrial applications, their use at the present time is limited by several factors, such as lack of cost-effective systems or processes for producing sufficient enzyme, heterogeneity of available preparations, and absence of lipases with properties required for certain applications (130). As with proteases, protein engineering can be applied to lipases to target numerous specific characteristics. Alteration of amino acid sequences will result in variants with modified specific activity, increased $k_{cat}$, altered pH and thermal activity profiles, increased stability (with respect to temperature, pH, and chemical agents such as oxidants and proteases), and show altered pI, surface hydrophobicity, and substrate specificity (160,161). Currently, lipase genes from fungal sources (e.g., G. candidum and C. rugosa) are being cloned and subjected to site-directed mutation to gain insight into structure–activity relationships, mainly with respect to selectivity, on which to base protein engineering strategies. Despite the enormous progress that has been made in this regard, the molecular basis for selectivity is still not well understood.

REFERENCES


I. INTRODUCTION

The development of methods to improve the nutritional and functional properties of fats and oils is of great interest to food processors. The molecular weight, unsaturation, and positional distribution of fatty acid residues on the glycerol backbone of triacylglycerols are the principal factors determining the physical properties of fats and oils (1,2). Chemical interesterification produces a complete positional randomization of acyl groups in triacylglycerols. It is used in the manufacture of shortenings, margarines, and spreads to improve their textural properties, modify melting behavior, and enhance stability (3,4). Interest in interesterification from a nutritional and functional standpoint is increasing since it can be used to produce margarines with no trans unsaturated fatty acids, synthesize cocoa butter substitutes and improve the nutritional quality of some fats and oils (5). Recently, research efforts have been directed to substituting some chemical interesterification applications with enzymatic interesterification because of the inherent advantages associated with the enzymatic process. Enzymatic reactions are more specific, require less severe reaction conditions, and produce less waste. Also, when immobilized, enzymes can be reused, thereby making them economically attractive (6). Interesterification, whether chemical or enzymatic, is the exchange of acyl groups between an ester and an acid (acidolysis), an ester and an alcohol (alcoholysis), an ester and an ester (transesterification) (7).

The major components of fats and oils are triacylglycerols, the composition of which is specific to the origin of each fat or oil. The physical properties of various fats and oils are different because of the structure and distribution of fatty acids in the triacylglycerols (8). In natural fats, acyl groups are distributed in a nonrandom fashion. During chemical or enzymatic interesterification, acyl groups are redistrib-
Enzymatic interesterification is accomplished using lipases, which are enzymes obtained predominantly from bacterial yeast, and fungal sources. Extracellular microbial lipases are produced by microorganisms and released into their growth environment to digest lipid materials (9). Lipases are defined as glycerol ester hydrolases (EC 3.1.1.3) because they catalyze the hydrolysis of carboxyl ester bonds in acylglycerols. Depending on the degree of hydrolysis, free fatty acids, monoacylglycerols, diacylglycerols, and glycerol are produced. Lipases are differentiated from esterases in that they act only on insoluble substrates. Long chain triacylglycerols, the natural substrates of lipases, are insoluble in water, forming aggregates or dispersions in aqueous media. Lipases have a high affinity for hydrophobic surfaces and can be completely adsorbed from aqueous solution by emulsified long chain triacylglycerols (11). In the presence of excess water, lipases catalyze the hydrolysis of long chain triacylglycerols, but under water-limiting conditions, the reverse reaction, ester synthesis, can be achieved (8,12). Enzymatic interesterification systems are composed of a continuous water-immiscible phase, containing the lipid substrate, and an aqueous phase containing the lipase. Lipase-catalyzed interesterifications have been extensively studied in systems using organic solvents. However, if such a process is to be used in the food industry, solvent-free systems must be developed. Hence, the emphasis of this chapter will be on enzymatic interesterification performed in solvent-free systems.

A. Transesterification

As previously defined, transesterification is the exchange of acyl groups between two esters, namely, two triacylglycerols (Fig. 1). Transesterification is used predominantly to alter the physical properties of individual fats and oils or fat–oil blends by altering the positional distribution of fatty acids in the triacylglycerols. Transesterification of butter using a nonspecific lipase has been reported to improve the plasticity of the fat (13). Kalo et al. (14) found that transesterification of butterfat with a positionally nonspecific lipase at 40°C increased the level of saturated C48 to C54 triacylglycerols, monoene C38 and C46 to C52 triacylglycerols, and diene C40 to C54 triacylglycerols. These authors also found that the diacylglycerol content increased by 45% whereas the free fatty acid content doubled. Overall, lipase-catalyzed tran-

![Figure 1](image-url)  
**Figure 1** Lipase-catalyzed transesterification between two different triacylglycerols.
esterification of butterfat at 40°C produced an increase in the solid fat content below 15°C and a decrease in the solid fat content above 15°C (Fig. 2).

In another study, lipase-catalyzed transesterification of butter increased the relative proportion of C36 and C40 to C48 saturated triacylglycerols, as well as triunsaturated triacylglycerols (15). The resulting product had a 114% greater solid fat content at 20°C than the starting butter, with the solid fat content increasing from 22% to 46%. In general, lipase-catalyzed transesterification produces fat with a slightly lower solid fat content compared with chemical interesterification. This is attributed to contamination by monoacylglycerols, diacylglycerols, and free fatty acids, which are produced in the early stages of transesterification (8,13). Kalo et al. (13) compared lipase-catalyzed transesterification to chemical interesterification of butter. He found that the solid fat content of butter increased from 41.2% to 42.2% at 20°C using lipase-catalyzed transesterification, whereas chemical interesterification produced butter with a solid fat content of 57.8% at 20°C. Transesterification has also been used to improve the textural properties of tallow and rapeseed oil mixtures as well as in the development of cocoa butter equivalents (16,17). Forsell et al. (18) found that transesterification of a tallow and rapeseed oil blend decreased the solid fat content and melting point. The extent of melting point reduction was dependent on the mass fraction of the two lipid components. With a mass fraction of tallow to rapeseed oil of 0.8, the melting point was reduced by 6°C, whereas a mass fraction of 0.5 produced a 12°C decrease in melting point. A decrease in the solid fat content has also been observed upon transesterification between palm oil and canola oil, due to a decrease in the level of triunsaturated triacylglycerols (19).

The attractiveness of cocoa butter to the chocolate and confectionary industry is based on the limited diversity of triacylglycerols in this fat, which gives it a unique, narrow melting range of 29–43°C. Chocolate can contain 30% cocoa butter, meaning that this fat determines the crystallization and melting properties of the chocolate. At 26°C, cocoa butter is hard and brittle, but when eaten it melts completely in the mouth with a smooth, cool sensation. The major triacylglycerols in cocoa butter

![Figure 2](image.png)

**Figure 2** Solid fat content versus temperature profiles for native and enzymatically interesterified butterfat in the absence of solvent using lipase from *Pseudomonas fluorescens*. Nontransesterified butterfat, (○); transesterified butterfat, (●). (From Ref. 13.)
are 1-palmitoyl-2-oleoyl-3-stearoylgllycerol (POS), 1,3-dipalmitoyl-2-oleoylglycerol (POP), and 1,3-distearoyl-2-oleoylglycerol (SOS) with levels of 41–52%, 16%, and 18–27%, respectively (8,17). The main disadvantage of using cocoa butter in chocolate and confections is its high cost. A cocoa butter equivalent can be made from inexpensive fats and oils by interesterification. By transesterifying fully hydrogenated cottonseed and olive oil, Chang et al. (17) were able to produce a cocoa butter substitute with similar POS levels and slightly higher SOS levels than those found in cocoa butter. The melting range of the transesterified product was 29–49°C, compared with 29–43°C for cocoa butter. In order to remove the desired triacylglycerol product from the other triacylglycerols, trisaturated triacylglycerols were removed by crystallization in acetone. High-oleic sunflower oil and palm oil fraction have also been transesterified to obtain cocoa butter substitutes (5).

B. Acidolysis

Acidolysis, the transfer of an acyl group between an acid and an ester, is an effective means of incorporating novel free fatty acids into triacylglycerols (Fig. 3). Acidolysis has been used to incorporate free acid or ethyl ester forms of eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) into vegetable and fish oils to improve their nutritional properties. The nutritional benefits of consuming polyunsaturated fatty acids, such as EPA and DHA, derived from fish oils have been proven. When consumed, EPA reduces the risk of cardiovascular disease by reducing the tendency to form blood clots, whereas DHA consumption is required for proper nervous system and visual functions, due to its accumulation in the brain and retina (20,21). Concentrations of EPA and DHA in fish oils to levels approaching 30% can be achieved using molecular distillation, winterization, and solvent crystallization. However, performing an acidolysis reaction between cod liver oil and free EPA and DHA, Yamane et al. (22) were able to increase the EPA content in the oil from 8.6% to 25% and the DHA content from 12.7% to 40% using immobilized lipase from \textit{Mucor miehei}. Using ethyl esters of EPA, fish oil has been enriched by interesterification to contain 40% EPA and 25% DHA (wt%) (23). During acidolysis in a fixed bed reactor, Yamane et al. (24) increased the polyunsaturated fatty acid (PUFA) content of cod liver oil by reducing the temperature to between −10°C and −20°C in the product reservoir. This led to crystallization and removal of more saturated fatty acids present in the fish oil. Lipases with strong specificities against EPA or DHA have also been used to enrich their content in fish oils (25). Future developments in lipase-catalyzed interesterification using EPA and DHA is directed to improving the nutritional quality of vegetable oils by enrichment with these fish oil–derived fatty acids. Acidolysis has also been used by Oba and Witholt (26) to incorporate oleic acid into milk fat. This process led to an increase in the level of unsaturated fatty acids in butter without losses in the characteristic flavor of butter. Acidolysis of milk fat with oleic acid was
also found to decrease the crystallization temperature and lower the melting range of the milk lipids.

Along with the enrichment of oils, acidolysis using EPA and DHA has also been useful in the synthesis of structured lipids. Structured lipids are composed of medium chain and long chain fatty acids, which meet the nutritional needs of hospital patients and those with special dietary needs. When consumed, medium chain fatty acids, such as capric and caproic acid, are not incorporated into chylomicrons and are therefore not likely to be stored, but will be used for energy. They are readily oxidized in the liver and constitute a highly concentrated source of energy for premature babies and patients with fat malabsorption disease. Medium chain fatty acids also possess a nutritional advantage compared with other fatty acids in that they are non-tumor-producing forms of fat (27). Long chain fatty acids are also required by the body, especially in the form of PUFAs in the form of omega-3 and omega-6 fatty acids, which have been associated with reduced risk of platelet aggregation and cardiovascular disease, and the lowering of cholesterol (27,28). When polyunsaturated fatty acids are present in the sn-2 position and medium chain fatty acids are present in the sn-1,3 positions, they are rapidly hydrolyzed by pancreatic lipase, absorbed and oxidized for energy, whereas essential PUFAs are absorbed as 2-monooacylglycerols. Therefore, structuring triacylglycerols with medium chain fatty acids and PUFAs can dramatically improve the nutritional properties of triacylglycerols (29). Producing a triacylglycerol rich in EPA or DHA at the sn-2 position, with medium chain fatty acids in the sn-1 or sn-3 positions, would provide maximal benefit, especially for intravenous use in hospitals (30). Structured lipids that are reduced in caloric content have also been developed by esterifying long chain monoacylglycerols containing behenic acid with capric acid. The produced triacylglycerols contain half the calories relative to natural triacylglycerols due to the incomplete absorption of behenic acid during digestion (31).

Acidolysis is also a common method for production of cocoa butter substitutes. The most common method is acidolysis of palm oil midfraction, which contains predominantly 1,3-dipalmitoyl-oleoyl-glycerol with stearic acid to increase the level of stearate in the lipid (32). Chong et al. (33) also incorporated stearic acid into palmolein to produce 25% cocoa butter-like triacylglycerols.

C. Alcoholysis

As previously mentioned, alcoholysis is the esterification reaction between an alcohol and an ester (Fig. 4). Alcoholysis has been used in the production of methyl esters from esterification of triacylglycerols and methanol with yields of up to 53% (34). During alcoholysis, hydrolysis of triacylglycerols to produce diacylglycerols and monoacylglycerols can occur, in some cases reaching levels as high as 11%, although the presence of small amounts of alcohol can inhibit hydrolysis. The main use of alcoholysis is in the performance of glycerolysis reactions.

![Figure 4](image)

**Figure 4** Lipase-catalyzed alcoholysis reaction between an acylglycerol and an alcohol.
Glycerolysis is the exchange of acyl groups between glycerol and a triacylglycerol to produce monoacylglycerols, diacylglycerols, and triacylglycerols. There are several ways to produce monoacylglycerols, which are of great importance in the food industry as surface-active agents and emulsifiers. Monoacylglycerols can be produced by ester exchange between triacylglycerols and glycerols, or by free fatty acids and glycerol, although only the former reaction is termed glycerolysis (Fig. 5). Glycerolysis is usually performed using nonspecific lipases, giving a wide range of reaction products (Fig. 6).

High yields in lipase-catalyzed monoacylglycerol synthesis are achieved by temperature-induced crystallization of newly formed monoacylglycerols from the reaction mixture. This pushes the equilibrium of the reaction toward increased monoacylglycerol production. Glycerolysis of lipids containing saturated monoacylglycerols in the reaction product mixture, since they crystallize at lower temperatures than unsaturated monoacylglycerols (35). Pseudomonas fluorescens and Chromobacterium viscosum have been shown to have high glycerolysis activity (36). In glycerolysis reactions, \( T_c \) is defined as the critical temperature below which monoacylglycerols formed by glycerolysis crystallize out of the reaction mixture. Removal of monoacylglycerols from the reaction mixture pushes the equilibrium of the reaction toward increased monoacylglycerol production. Vegetable oils with low melting points due to the presence of long chain unsaturated fatty acids have a much lower

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**Figure 5** Lipase-catalyzed glycerolysis reaction between glycerol and a triacylglycerol to produce monoacylglycerols.

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**Figure 6** Products of a nonspecific lipase-catalyzed glycerolysis reaction between glycerol and 1,3-dipalmitoyl-2-oleoyl-glycerol.
than animal fats. The \( T_c \) for vegetable oils ranges from 5°C to 10°C, whereas it is between 30°C and 46°C for animal fats. By reducing the temperature below \( T_c \), yields of monoacylglycerols can be increased from 30% up to yields as high as 90% (35,36). Water content can also have an effect on glycerolysis since the reaction is an esterification. McNeil et al. (36) found that increasing the water content from 0.5% to 5.7% increased the production of monoacylglycerols, whereas higher levels of water did not increase the rate of reaction further. The main problem with lipase-catalyzed glycerolysis is the long reaction time in the order of 4–5 days required to produce high yields (36).

II. LIPASES

A. Three-Dimensional Structure

While lipases can be derived from animal, bacterial, and fungal sources, they all tend to have similar three-dimensional structures. In the period from 1990 to 1995, crystallographers solved the high resolution structures of 11 different lipases and esterases including 4 fungal lipases, 1 bacterial lipase, and human pancreatic lipase (12). Comparison of the amino acid sequences has shown large differences between most lipases, yet all have been found to fold in similar ways and have similar catalytic sites. The characteristic patterns found in all lipases studied so far have included \( \alpha/\beta \) structures with a mixed central \( \beta \) sheet containing the catalytic residues. In general, a lipase is a polypeptide chain folded into two domains: the C-terminal domain and the N-terminal domain. The N-terminal domain contains the active site with a hydrophobic tunnel from the catalytic serine to the surface that can accommodate a long fatty acid chain.

In solution, a helical segment covers the active site of lipase, but in the presence of lipids or organic solvent, there is a conformational change in which the lid opens, exposing the hydrophobic core containing the active site. The structure of the lid differs for lipases in the number and position of the surface loops. For example, human pancreatic lipase has one \( \alpha \) helix (residues 237–261) in the loop covering the active site pocket (37,38). The fact that the \( \alpha \) helix in the lid is amphipathic is very important in terms of the ability of the lipase to bind to lipid at the interface. If the amphiphilic properties of the loop are reduced, the activity of the enzyme is decreased (39). The outside of the loop is relatively hydrophilic whereas the side facing the catalytic site is hydrophobic. Upon association with the interface, the lid folds back, revealing its hydrophobic side which leads to increased interactions with the lipid at the interface (40). The substrate can then enter the hydrophobic tunnel containing the active site.

B. The Active Site

Koshland’s modern induced fit hypothesis states that the active site does not have to be a preexisting rigid cavity but instead can be a precise spacial arrangement of several amino acid residues that are held in the correct orientation by the other amino acids in the enzyme molecule (41). The main component of the catalytic site is an \( \alpha/\beta \)-hydrolase fold that contains a core of predominantly parallel \( \beta \) sheets surrounded by \( \alpha \) helices. The folding determines the positioning of the catalytic triad composed of serine, histidine, and either glutamic acid or aspartic acid along with several...
oxyanion-stabilizing residues. The nucleophilic serine rests between a $\beta$ strand and an $\alpha$ helix, whereas histidine and aspartic acid or glutamic acid rest on one side of the serine (12).

The importance of the serine residue for the catalytic activity of lipase has been demonstrated using site-directed mutagenesis. Substitution of Ser153 in human pancreatic lipase produces a drastic decrease in the catalytic activity of the enzyme, but has no effect on the ability of the enzyme to bind to micelles. As well, the presence of a highly hydrophobic sequence of amino acid residues has been verified in the vicinity of the active site, which is important in the interaction of the enzyme with the interface (42). The chemical properties of the groups within the catalytic triad are consistent with a hydrophobic environment (11). The process of opening the lid covering the active site causes the oxyanion hole to move into proper positioning for interaction with the substrate. For example, lipase for *Rhizomucor miehei* has a serine side chain at position 82 that assumes a favorable conformation for oxyanion interactions only after the lid has moved away from the active site (43). During binding of the substrate with the enzyme, an ester binds in the active site, so that the alcohol portion of the substrate rests on a floor formed by the end of the $\beta$ strand while the acyl chain arranges itself in the hydrophobic pocket and tunnel region (42) (Fig. 7).

In lipase from *Mucor miehei*, the substrate binding region is seven carbons long. When longer chains are encountered, the rest of the carbons in the chain hang outside the hydrophobic tunnel (44). When the lipase approaches the interface and the lid is folded back, an oxyanion-stabilizing residue is placed in proper orientation (12). During hydrolysis the tetrahedral intermediate is stabilized by hydrogen bonds with backbone amide groups of oxyanion-stabilizing residues. One stabilizing residue

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**Figure 7** Crystal structure and location of catalytic residues of the active site of *Candida rugosa* lipase. (Adapted from Ref. 37.)
in the oxyanion hole is the amino acid following the catalytic serine, whereas the other one comes from a separate loop (37).

C. Activation by Interfaces

As previously stated, an advantage of enzyme-catalyzed interesterification in comparison with chemical methods is that it can operate effectively under relatively mild conditions. Enzyme-catalyzed reactions can increase the rate of a reaction by $10^6$–$10^{15}$ times even at 25°C (41). The kinetics of lipase-catalyzed interesterification can get complicated due to the many factors that can affect the reaction. Activation by interfaces as well as participation of multiple substrates in the interesterification reaction must all be considered when describing the action of lipases at interfaces.

The natural substrates of lipases, long chain triacylglycerols, are uncharged and insoluble in water and as such form two phases in aqueous solutions. The property of being active at lipid–water interfaces is unique to lipases. At low concentrations of lipids, termed monomeric solutions, the lipids are dissolved in aqueous phase. The maximal concentration of monomers in aqueous solution is the solubility limit or critical micelle concentration, after which triacylglycerols form emulsions. For example, the critical micelle concentration for triacetin in aqueous solution is 0.33 M, whereas for long chain triacylglycerols, it can be as low as 1.0 μM (12,45). It has been shown that lipases display almost no activity toward monomeric solutions of lipids, whereas the lipids are dissolved and do not form interfaces. Once the level of lipids exceeds the critical micellar concentration, the reaction rate increases dramatically, by a factor of $10^3$–$10^4$ in some cases depending on the quality of the interface (Fig. 8). Lipases have been found to act at several interfaces, including emulsions, bilayers, and micelles (46). Action of lipases at the lipid–water interface is believed to follow two successive equilibria involving penetration of lipase into the interface, followed by the formation of the enzyme substrate complex (Fig. 9).

Initially, the enzyme penetrates the interface and undergoes a conformational change, folding back the lid and thereby increasing the hydrophobic surface area of

![Figure 8](Image)

**Figure 8** Comparison of the effect of substrate concentration on lipase and esterase activity at monomeric and saturation levels (beyond vertical dashed lines). (From Ref. 45.)
the lipase making contact with the interface. The enzyme adsorbs to the interface following a Langmuir adsorption isotherm. Once adsorption has taken place, the enzyme is in its catalytically active form, meaning that interfacial activation has taken place. The lipid substrate can then fit into the active site and be transformed into product. The product is believed to be water-soluble and leave the interface rapidly by diffusion into the surrounding solution. Several mechanisms have been proposed to explain interfacial activation of lipases. The first theory relates interfacial activation to a conformational change of the enzyme, where the lid moves to make the active site available to substrate molecules at the interface. The second theory points to changes in the concentration and organization of substrate molecules at the interface to cause activation of the lipase. In the presence of a non-substrate lipid interface, a lipase will not be active, but once the concentration of substrate in the interface exceeds that of non-substrate lipids to become the continuous phase, lipase activity increases. There are several other theories as to why lipase activity is increased at an interface. One theory states that the higher substrate concentration at the interface produces more frequent collisions between the lipase and substrate than in monomeric solutions. Other theories involve decreased energy of activation induced by substrate aggregation, reduced hydration of the substrate, and progressive lipid-induced lipase aggregation at the interface (46).

In considering the action of lipases at interfaces, several factors have to be considered, including the reversibility of adsorption, the possibility of inactivation, and the quality of the interface. In general, lipases are considered to be reversibly adsorbed at interfaces, since by increasing surface pressure, lipases have been found to desorb from the interface (46). The quality of the interface can affect the activity of lipases. Any factor that affects the affinity of the enzyme for the interface as well as packing and orientation of the molecules at the interface can affect activity (11).

D. The Problem of Substrate Concentration

Since long chain triacylglycerols are insoluble in water and form aggregates, lipase-catalyzed interesterification cannot be strictly governed by the Henri–Michaelis rule relating the rate of the reaction to the molar concentration of substrate in solution. In interesterification reactions, the insoluble substrate is in large excess as the continuous solvent phase, making it difficult to define its concentration in the reaction mixture. Since the substrate is insoluble, only the concentration of the substrate present at the interface, which is available to the lipase, is considered. Lipase activity is controlled by the concentration of micellar substrates at the interface and is independent of the molar concentration of the substrate (47). In contrast, esterases in acting only on water-soluble substrates, have a Michaelis–Menten dependence on substrate concentration (42,48). The dependence of lipase activity on the surface area...
of the interface as a measure of substrate concentration was proven by Bezonana and Desnuelle (49), who measured the rates of hydrolysis in coarse and fine emulsions (Fig. 10). When the rate of the reaction for the two emulsions was plotted as a function of substrate mass, the same maximal rates of hydrolysis were obtained; however, there was a difference in the values for $K_m$. In contrast, when initial velocities were plotted as a function of interfacial area, the values of $K_m$ and $V_{max}$ were constant for both fine and coarse emulsions, indicating that the concentration of substrate at the interface (mol/m$^3$) directly determines the rate of the reaction (11,12,46,50). Reaction rates have also been shown to be a function of the emulsion concentration. If a stock emulsion is diluted to different concentrations, a progress curve of rate versus concentration will be obtained, which upon plotting gives a straight line in the Lineweaver–Burke plot. A relative $K_m$ can be obtained from this plot, but in order to obtain the absolute value for $K_m$, the area of the interface must be known (11). It is very difficult to obtain an accurate assessment of the interfacial area due to several factors. In free enzyme solutions, the size distribution of emulsion droplets and the degree of adsorption of enzyme to the interface must be known. It is difficult to estimate the surface area of the interface due to size heterogeneity and the possibility of coalescence of emulsion droplets. With immobilized enzyme, the size distribution and surface area of support particles and pores must be determined, as well as the degree of loading of the lipase (47). Due to the difficulty in measuring these factors accurately, only relative $K_m$ values are determined.

E. Kinetics and Mechanism of Action

Interesterification is a multisubstrate reaction, with the main substrates being glycerides, fatty acids, and water. This reaction can be considered a special case of chemical group transfer, involving sequential hydrolysis and esterification reactions (51). Lipase-catalyzed interesterification follows a Ping-Pong Bi-Bi reaction for multisubstrate reactions (50,51). The actual mechanism of acylation and deacylation of
the glyceride in the active site is shown in Figure 11. During acylation, a covalent acyl–enzyme complex is formed by nucleophilic attack of the active site serine on the carbonyl carbon of the substrate. The serine is made a stronger nucleophile by the presence of histidine and aspartic acid residues. The histidine imidazole ring becomes protonated and positively charged, stabilized by the negative charge of the active site aspartic acid or glutamic acid residues. A tetrahedral intermediate is subsequently formed, stabilized by two hydrogen bonds formed with oxyanion-stabilizing residues (12). A break in the carbon–oxygen bond of the ester causes release of the alcohol. During the reaction, the acylglycerol is associated with the catalytic triad through covalent bonds. Histidine hydrogen bonds with both serine and the oxygen of the leaving alcohol. Nucleophilic attack by water or an alcohol causes the addition of a hydroxyl group to the carbonyl carbon, producing a tetrahedral intermediate, which will rearrange, releasing the altered acylglycerol and regenerating the active site serine (42,52).

The first stage of interesterification involves hydrolysis of triacylglycerols with consumption of water to produce diacylglycerols, monoacylglycerols, and free fatty acids. Accumulation of hydrolysis products will continue during interesterification until an equilibrium is established (51). Since lipases are involved in multisubstrate, multiproduct reactions, more complex kinetic mechanisms are required. Interestefication involves acylation and deacylation reactions, either of which can be the rate-limiting step (50,53). The basic mechanism for a Ping-Pong Bi-Bi reaction using multiple substrates is shown in Figure 12.

Under steady-state conditions,

\[
\frac{v}{V_{\text{max}}} = \frac{[AX][BX]}{K_{m_{AX}}[AX] + K_{m_{AX}}[B] + [A][B]}
\]

where AX is the first substrate and BX is the second substrate (41). It is difficult to study the kinetics of Ping-Pong Bi-Bi mechanisms due to the presence of two substrates. In order to study the kinetics, one substrate concentration is usually held constant while the other one is altered. In the case of lipase-catalyzed interesterification under aqueous conditions, there is the additional difficulty that the lipid substrate is also the reaction medium, which is in excess compared with other components. Even with measurable amounts of lipid substrate, it is difficult to develop rate equations since all species involved have to be considered (50).

F. Specificity

The main advantage of lipases that differentiates enzymatic interesterification from chemical interesterification is their specificity. The fatty acid specificity of lipases has been exploited to produce structured lipids for medical foods and to enrich lipids with specific fatty acids to improve the nutritional properties of fats and oils. There are three main types of lipase specificity: positional, substrate, and stereo. Positional and fatty acid specificity are usually determined by partial hydrolysis of synthetic triacylglycerols and separation by thin-layer chromatography with subsequent extraction and analysis of the products. Other methods include conversion of the fatty acids produced during hydrolysis to methyl esters for gas chromatographic analysis (54).
Figure 11  Catalytic mechanism for lipase-catalyzed interesterification, showing the catalytic site containing Asp/Glu, His, and Ser residues. (Adapted from Ref. 52.)
1. Nonspecific Lipases

Certain lipases show no positional or fatty acid specificity during interesterification. Interesterification with these lipases after extended reaction times gives complete randomization of all fatty acids in all positions and gives the same products as chemical interesterification (Fig. 13). Examples of nonspecific lipases include lipases derived from *Candida cylindraceae*, *Corynebacterium acnes*, and *Staphylococcus aureus* (9,10).

2. Positional Specificity

Positional specificity, i.e., specificity toward ester bonds in positions sn-1,3 of the triacylglycerol, results from an inability of lipases to act on position sn-2 on the triacylglycerol, due to steric hindrance (Fig. 14). Steric hindrance prevents the fatty acid in position sn-2 from entering the active site (9,55). An interesterification reaction using a 1,3-specific lipase will initially produce a mixture of triacylglycerols, 1,2- and 2,3-diacylglycerols, and free fatty acids (55). After prolonged reaction periods, acyl migration can occur, with the formation of 1,3-diacylglycerols, which allows some randomization of the fatty acids existing at the middle position of the triacylglycerols. In comparison with chemical interesterification, 1,3-specific lipase-catalyzed interesterification of oils with a high degree of unsaturation in the sn-2 position of the triacylglycerols will decrease the saturated to unsaturated fatty acid level (56). Lipases that are 1,3-specific include those from *Aspergillus niger*, *Mucor miehei*, *Rhizopus arrhizus*, and *Rhizopus delemar* (9). The specificity of individual lipases can change due to microenvironmental effects on the reactivity of functional groups or substrate molecules (57). For example, lipase from *Pseudomonas fragi* is known to be 1,3-specific but has also produced random interesterification, possibly due to a microemulsion environment. As of yet, lipases that are specific toward fatty acids in the sn-2 position have been difficult to identify. Under aqueous conditions, one such lipase from *Candida parapsilosis* hydrolyzes the sn-2 position more rapidly than either of the sn-1 and sn-3 positions, and is also specific toward long chain polyunsaturated fatty acids (58).

The differences in the nutrition of chemically interesterified fats and oils compared to enzymatically interesterified samples can be linked to the positional specificity exhibited by some lipases. In fish oils and some vegetables oils that contain high degrees of essential PUFAs, these fatty acids are usually found in greater quantities in the sn-2 position. In the intestines, 2-monoacylglycerols are more easily absorbed than sn-1 or sn-3 monoacylglycerols. Using a 1,3-specific lipase, the fatty acids are acylated more readily in the sn-2 position, which may allow for the more complete absorption of essential fatty acids.
acid composition of positions 1 and 3 can be changed to meet the targeted structural requirements while retaining the nutritionally beneficial essential fatty acids in position 2. Using random chemical interesterification, retention and improvement in beneficial fatty acid content cannot be accomplished due to the complete randomization of the fatty acids in the triacylglycerols (59).

3. Stereospecificity

In triacylglycerols, the $sn$-1 and $sn$-3 positions are sterically distinct. Very few lipases differentiate between the two primary esters at the $sn$-1 and $sn$-3 positions, but when they do, the lipases possess stereospecificity. In reactions where the lipase is stereospecific, positions 1 and 3 are hydrolyzed at different rates. Stereospecificity is determined by the source of the lipase and the acyl groups, and can also depend on the lipid density at the interface, where an increase in substrate concentration can decrease specificity due to steric hindrance. Differences in chain length can also affect the specificity of the lipase (12). Lipase from *Pseudomonas* species and porcine pancreatic lipase have shown stereoselectivity when certain acyl groups are hydrolyzed (60).

4. Fatty Acid Specificity

Many lipases are specific toward particular fatty acid substrates. Most lipases from microbial sources show little fatty acid specificity, with the exception of lipase from *Geotrichum candidum*, which is specific toward long chain fatty acids containing cis-9 double bonds (9). Lipases can also demonstrate fatty acid chain length specificity, with some being specific toward long chain fatty acids and others being specific toward medium chain and short chain fatty acids. For example, porcine pan-
creatic lipase is specific toward shorter chain fatty acids, while lipase from *Penicillium cyclopium* is specific toward long chain fatty acids. As well, lipases from *Aspergillus niger* and *Aspergillus delemar* are specific toward both medium chain and short chain fatty acids (11,61). Other lipases have been found to be specific toward fatty acids of varying lengths. Marangoni (62) found that in the hydrolysis of butter oil, lipase from *Candida rugosa* showed specificity toward butyric acid compared to *Pseudomonas fluorescens* lipase. With interesterification reactions in organic media, lipases can also be specific toward certain alcohol species. A large group of lipases from sources such as *Candida cylindraceae*, *Mucor miehei*, and *Rhizopus arrhizus* have been found to be strongly specific against fatty acids containing the first double bond from the carboxyl end at an even-numbered carbon, such as cis-4, cis-6, and cis-8, resulting in slower esterification of these fatty acids in comparison with other unsaturated and saturated fatty acids. Fatty acid specificity by certain lipases can be used in the production of short chain fatty acids for use as dairy flavors and in the concentration of EPA and DHA in fish oils by lipases with lower activity toward these fatty acids.

### III. REACTION SYSTEMS

#### A. Enzymatic Interesterification in Microaqueous Organic Solvent Systems

Since the main substrates of lipases are long chain triacylglycerols, which are insoluble in water, many experiments have been conducted in the presence of organic solvents. Organic solvents allow the fat or oil to be solubilized and convert two-phase systems to one-phase systems (63). Stability can be improved by covalent attachment of polyethylene glycol to free amino groups of the lipase, giving lipases amphiphilic properties and allowing their dissolution in organic solvents (64). It has been reported that the thermal stability of lipases can be improved in microaqueous organic solvent systems since the lack of water prevents unfolding of the lipase at high temperatures (65). Elliott and Parkin (65) found that porcine pancreatic lipase had optimal activity at 50°C in an emulsion, whereas the optimum increased to 70°C in a microaqueous organic solvent system using hexane. Lipase activity in organic solvents depends on the nature and concentration of the substrate and source of enzyme (63). The specific organic solvent used can dramatically affect the activity of the lipase (66). Lipases are more active in *n*-hexane and isooctane than other solvents, such as toluene, ethyl acetate, and acetylnitrile (28,44). The polarity of solvents can be described by $P$, the partition coefficient of a solvent between water and octanol. This is an indication of the hydrophobicity of the solvent. No lipase activity is observed in solvents with a value for log $P$ less than 2 (67,68). The hydrophobicity of the solvent can also affect the degree of acyl migration during interesterification using a 1,3-specific lipase. Hexane tends to promote acyl migration due to the low solubility of free fatty acids and partial glycerides in hexane, which forces them into the microaqueous region around the lipase, providing optimum conditions for acyl migration. In contrast, the use of diethyl ether, in which free fatty acids and partial glycerides are more soluble, removes the products from the microaqueous environment and reduces the risk of acyl migration (6). Since the choice of organic solvents based on minimization of acyl migration may conflict with max-
imization of interesterification, acyl migration is usually minimized simply by reducing reaction times. Lipases can be made more active and soluble in organic solvent systems by attachment of an amphiphilic group such as polyethylene glycol (PEG). PEG reacts with the N-terminal or lysine amino groups, rendering the lipase more soluble in organic solvents (69). The activity of lipases in organic solvent depends on the solubility of the solvent in water. Lipases are only active in water-immiscible solvents, since water-miscible organic solvents extract the water of hydration layer from the vicinity of the enzyme, thereby inactivating them (44). Since the success of an interesterification reaction depends on the concentration of water in the system, the hydration state of the lipase plays a key role because a minimal amount of water is needed to maintain the enzyme in its active form. The use of hydrophobic solvents limits the flexibility of the enzyme, preventing it from assuming its most active conformation. Therefore, if organic solvents are used, the enzyme must be in its active conformation prior to addition of the organic solvent. This can be accomplished by exposing the enzyme to an inhibitor or substrate, then drying it in its active conformation (12,70). The advantage of using organic solvents in lipase-catalyzed interesterification reactions is that the water content can be carefully controlled. A water content higher than 1% can produce high degrees of hydrolysis, whereas water levels lower than 0.01% can prevent full hydration of the lipase and reduce the initial rate of hydrolysis (1). Therefore, water levels between these two extremes are necessary to maximize the effectiveness of enzymatic interesterification in organic solvents. In microaqueous organic solvent systems, the effect of pH on lipase activity is complex because water levels are so low. It has been proven that enzymes in organic solvent systems have a memory of the pH of the last aqueous environment in which they were. Elliott and Parkin (65) found that porcine pancreatic lipase has an optimum activity in hexane after being exposed to pH values between 6.5 and 7.0. At pH 8.5, the decrease in activity was attributed to a change in the ionization state of the histidine in the active site.

A common form of organic solvent system used in lipase-catalyzed interesterification is that of reverse micelles. Reverse micelles, or microemulsions, are defined as nanometer-sized water droplets dispersed in organic media with surfactants stabilizing the interface (29,71). A common surfactant used is an anionic double-tailed surfactant called sodium-bis(2-ethylhexyl)sulfosuccinate (AOT). Reverse micelles are used in interesterification reactions because they increase the interfacial area and improve the interaction between lipase substrate (29). As well, the use of microemulsions makes it possible to use polar and nonpolar reagents in the same reaction mixture (72). Reverse micelles can be formed by gently agitating a mixture of AOT, lipid substrate, organic solvent, and lipase dissolved in buffer until the solution becomes clear. The lipase is trapped in an aqueous medium in the core of the micelle, avoiding direct contact with the organic medium (61). Lecithin has been used to promote the formation of reverse micelles and to protect the lipase from nonpolar solvents (73,74). At ionic strengths higher than 1 M, activity is decreased due to decreased solubility and activity of the lipase. The water content required for microemulsion systems is dependent on the desired reaction, although some level of water is necessary to hydrate the enzyme. For example, Holmberg et al. (75) found that 0.5% water was the optimum for production of monoacylglycerols from palm oil in a microemulsion. The composition of the substrate can also affect the rate of interesterification in reverse micelles. Substrates with more amphiphilic properties
are better because they can partition to the interface. More polar substrates tend to stay in the water phase and interact less with the interface (76). The disadvantages of reverse micelle systems are that lipase activity is decreased rapidly, and the system can alter lipase specificity (73,76,77). Reverse micelles can also be used with immobilized lipases, where the reverse micelle is formed around the support and immobilized lipase. This method has been used with hexane to produce cocoa butter equivalents (73). Although they have been used in experimental form to produce triacylglycerols from diacylglycerols and oleic acid (78), as well as triacylglycerols suitable for use as cocoa butter substitutes (74), reverse micelles are not used in industrial enzymatic interesterification applications.

IV. IMMOBILIZATION

Immobilization of lipases has become increasingly popular for both hydrolysis and synthesis reactions. The advantages of immobilized enzyme systems compared to free enzyme systems include reusability, rapid termination of reactions, lowered cost, controlled product formation, and ease of separation of the enzyme from the reactants and products. In addition, immobilization of different lipases can affect their selectivity and chemical and physical properties. Immobilization also provides the possibility of achieving both purification of the lipase from an impure extract and immobilization simultaneously, with minimal inactivation of the lipase (79). Methods for immobilization of enzymes include chemical forms, such as covalent bonding, and physical forms, such as adsorption and entrapment in a gel matrix or microcapsules (7,80).

The easiest and most common type of immobilization used in interesterification reactions is adsorption, which involves contacting an aqueous solution of the lipase with an organic or inorganic surface-active adsorbent. The objective of immobilization is to maximize the level of enzyme loading per unit volume of support. The process of adsorption can be accomplished through ion exchange or through hydrophobic or hydrophilic interactions and van der Waals interactions (81). After a short period of mixing of the free enzyme and support, the immobilized enzyme is washed to remove any free enzyme that is left, after which the product is dried (79). The same adsorption process can be accomplished by precipitating an aqueous lipase solution onto the support using acetone, ethanol, or methanol, then drying as previously described (9,81). Although desorption can occur, most immobilized lipase preparations are stable in aqueous solutions for several weeks. The preparations are stable because as the lipase adsorbs to the support, it unfolds slightly, allowing several points of interaction between the lipase and support. In order for desorption to occur, simultaneous loss of interactions at all contact sites must occur, which is unlikely (82).

The degree of immobilization depends on several conditions, including pH, temperature, solvent type, ionic strength, and protein and adsorbent concentrations. The choice of carrier is dependent on its mechanical strength, loading capacity, cost, chemical durability, functionality, and hydrophobic or hydrophilic character (83). In general, lipases retain the highest degree of activity when immobilized on hydrophobic supports, where desorption of lipase from the support after immobilization is negligible, and improved activity has been attributed to increased concentrations of hydrophobic substrate at the interface (7,50). The disadvantages of using hydrophilic
supports include high losses of activity due to changes in conformation of the lipase, steric hindrance, and prevention of access of hydrophobic substrates (7). Common hydrophobic supports include polyethylene, polypropylene, styrene, and acrylic polymers, while hydrophilic supports include Duolite, Celite, silica gel, activated carbon, clay, and Sepharose (7). The effectiveness of the immobilization process is influenced by the internal structure of the support. If a support with narrow pores is used, most of the enzyme will be immobilized on the surface of the support, which prevents the occurrence of internal mass transfer limitations. If a support containing larger pore sizes is used, such as Spherosil DEA, with an average diameter of 1480 Å, some lipase will be immobilized inside the pores, which can prevent access of the substrate to some of the lipase. This is due to preferential filling of pores and crevices by the lipase during immobilization (84,85). The activity of lipases tends to decrease upon immobilization, with activity being reduced by 20–100% (79,81). The activity of an immobilized enzyme relative to the free form can be compared by an effectiveness value, which is defined as the activity of immobilized enzyme divided by the activity of an equal amount of free enzyme determined under the same operating conditions. The effectiveness value can be used as a guide to the degree of inactivation of the enzyme caused by immobilization. For values close to 1.0, very little enzyme activity has been lost upon immobilization, whereas values much lower than 1 indicate high degrees of enzyme inactivation (80).

The performance of an immobilized lipase can also be affected by handling and reaction conditions. Freeze drying of the immobilized enzyme before interesterification to substantially reduce the moisture content has been reported to dramatically improve activity. Molecular sieves can also be added to reaction systems to reduce the amount of water that accumulates during the reaction, which would in turn reduce the degree of hydrolysis (4). The main disadvantage associated with adsorption as an immobilization method is that changes in pH, ionic strength, or temperature can cause desorption of lipase that has been adsorbed by ion exchange. Lipases adsorbed through hydrophobic or hydrophilic interactions can be desorbed by changes in temperature or substrate concentration (79).

A. Factors Affecting Immobilized Lipase Activity

Immobilization can have an impact on the activity of lipases through steric, mass transfer, and electrostatic effects. During immobilization, the enzyme conformation can be affected and parts of the enzyme can be made inaccessible to the substrate due to steric hindrance.

1. Mass Transfer Effects

The kinetics of lipase-catalyzed interesterification can be affected by mass transfer limitations. The substrate must diffuse through the fluid boundary layer at the surface of the support into the pore structure of the support and react with the lipase. Once products have been released by the lipase, they must diffuse back out of the pore structure and away from the surface of the support. Mass transfer limitations fall into two categories: internal and external mass transfer. Internal mass transfer is the transport of substrate and product within the porous matrix of the support; it is affected by the size, depth, and smoothness of the pores. Internal mass transfer is diffusion-limited only. When the rate of diffusion inward is slower than the rate of conversion.
of substrate to product, the reaction is diffusion-limited, as there is not enough substrate available for the amount of enzyme present (86). A diffusion coefficient for internal mass transfer in immobilized enzyme systems compared to free enzyme systems is defined as:

\[
D_e = \frac{D \Psi}{\tau}
\]

where \(D_e\) is the effective diffusion coefficient inside the support particles, \(D\) is the diffusion coefficient in free solution, \(\Psi\) is the porosity of the particles, and \(\tau\) is the tortuosity factor, defined as the distance of the pathlength traveled by molecules between two points in a particle.

The effective diffusion coefficient varies inversely with the molecular weight of the substrate (80). Internal diffusional limitations can be recognized if the activity increases when the support particles are crushed, since crushing would decrease the length of the pathway that the substrate would have to travel to reach the enzyme. The Thiele modulus, \(\phi\), can be used to evaluate the extent of internal mass transfer limitations:

\[
\frac{1}{2} \frac{V_{\text{max}}}{K_D} = \frac{L}{L^2 K_D} = \frac{L}{L^2 K_D} = L
\]

where \(L\) is the half-thickness of the support particles. Internal mass transfer limitations can also be identified by measuring the initial velocity of the reaction at increasing enzyme concentrations. If the rate of the reaction remains constant at increasing enzyme concentrations (amount of enzyme per gram of support), the reaction is mass transfer–limited. If the rate of reaction increases linearly with increasing enzyme concentration, the reaction is kinetically limited. Internal diffusion limitations can be reduced by decreasing the support particle size, increasing pore size and smoothness, using low molecular weight substrates, and using high substrate concentrations (80). The difficulty with using smaller support particles in fixed bed reactors where internal mass transfer limitations are high is that it tends to increase the back pressure of the system (84).

External mass transfer limitations are the resistance to transport between the bulk solution and a poorly mixed fluid layer surrounding each support particle. External mass transfer can occur in packed bed and membrane reactors and is affected by both convection and diffusion (84). If the reaction is faster than the rate of diffusion of substrate to the surface or product from the surface, this can affect the availability of substrate for lipase catalysis. If inadequate substrate quantities reach the enzyme, the rate of reaction will be lower than that of free enzyme. An increasing external mass transfer coefficient can be identified during kinetic analysis by an increasing slope of a Lineweaver–Burk plot (87). In stirred reaction systems, external mass transfer limitations can be reduced in packed bed reactors by increasing the flow rate, reducing the viscosity of the substrate, and increasing substrate concentration (80). Changing the height-to-diameter ratio of a fixed bed reactor can also reduce external mass transfer limitation as it increases the linear velocity of the substrates.

2. The Nernst Layer and Diffusion Layer

Immobilized lipases are surrounded by two different layers, which can create differences in substrate concentration between them and the bulk phase. The Nernst layer
is a thin layer located directly next to the surface of the support. In the case of hydrophobic supports and hydrophobic substrates, such as triacylglycerols, the concentration of substrates in the Nernst layer is more concentrated than in the bulk solution since the hydrophobic substrate tends to partition toward the hydrophobic support material. Another layer surrounding the support particles is a diffusion or boundary layer. A concentration gradient is established between the diffusion layer and the bulk phase as substrate is converted to product by the lipase. The product concentration in the diffusion layer is higher than in the bulk phase as it must diffuse from the surface of the support into the bulk phase. Consequently, due to the higher product concentration in the diffusion layer, the substrate concentration is lower than in the bulk phase, producing concentration gradient with more substrate diffusing toward the support and immobilized lipase. Differences in substrate concentration between the Nernst layer and/or the boundary layer and the bulk phase can affect the determination of $K_m$ since substrate concentration will be measured in the bulk layer, which may not be the concentration of substrate closer to the lipase. With a lower substrate concentration at the support in comparison with the bulk phase, the apparent $K_m$ will appear higher and the activity will appear lower than its actual values. The opposite will occur with a higher substrate concentration at the interface.

A third factor that can affect the activity of immobilized lipase is electrostatic effects. If the support and substrate possess the same charge then they will experience repulsion, whereas if they have opposite charge they will be attracted. This factor can have an effect on the apparent $K_m$. As well, electrostatic effects can have an impact on other components in the reaction. For example, if the support was anionic, the local concentration of hydrogen ions would be higher in the vicinity of the immobilized lipase, which would cause a decrease in the pH around the enzyme.

Combining the electrostatic effects and the effect of the Nernst layer, the value of the apparent $K_m$ can be modified as follows (88):

$$K_{m}' = \left( K_m + \frac{x}{D} V_{max} \right) \frac{RT}{RT - \alpha z F V}$$

where $K_{m}'$ is the apparent $K_m$ of the lipase, $x$ is the thickness of Nernst layer, $R$ is the universal gas constant, $T$ is the absolute temperature, $\alpha$ is the valence of the substrate, $F$ is Faraday's constant, $V$ is the magnitude of the electric field around the enzyme support, and $D$ is the diffusion coefficient of the substrate.

If the thickness of the Nernst layer decreases, then the ratio $x/D$ would decrease and $K_{m}'$ would decrease, approaching $K_m$.

B. Stability of Immobilized Enzymes

The stability of immobilized enzymes depends on the method of immobilization and the susceptibility of the enzyme to inactivation. Inactivation can be caused by contaminants and changes in temperature, pH, and ionic strength. High shear, microbial contamination, fouling, and breakage of support particles have also been found to inactivate immobilized enzymes. Depending on the strength of the immobilization method, the enzyme can also be desorbed from the support. The stability of immobilized enzymes is evaluated by determining the half-life of the enzyme under the reaction conditions. In diffusion-limited systems, there is a linear decay in enzyme activity in time, as enzymes on the surface of the support are inactivated and the
substrate diffuses further into the pores to reach enzyme molecules that have not been inactivated. In systems free of diffusional limitations, enzyme inactivation follows a first-order decay:

\[
\ln \frac{N_0}{N} = \lambda t
\]

where \(N_0\) is the initial enzyme activity, \(N\) is the activity at time \(t\), and \(\lambda\) is the decay constant. Using \(\lambda\), the half-life of the immobilized lipase can be determined as follows:

\[
\text{half-life} = \frac{0.693}{\lambda}
\]

The half-lives of lipases in interesterification systems have been reported to range from 7 minutes to 7 months, with the large variability attributed to the source of lipases and different reaction conditions (50). As previously stated, the half-life of the immobilized enzyme can be used to determine the productivity of the system. In order to avoid losses in productivity as the activity of the immobilized lipase decreases, the temperature can be raised to increase the reaction rate or, in fixed bed reactor systems, the flow rate can be reduced (80). While these measures can improve the conversion rate, they can also increase the rate of enzyme inactivation in the case of temperature increases, or decrease the throughput in the case of reduced flow rate.

C. Immobilized Enzyme Kinetics

The previous discussion on the kinetics of lipase action was developed for soluble lipases acting on insoluble substrate, but assuming that diffusional and mass transfer effects are not rate-limiting, the same theories can be applied to immobilized lipases. When using immobilized lipases, the level of substrate in comparison with the level of enzyme must be considered. In general, there is a low average concentration of substrate in direct contact with the immobilized lipase due to high conversion rates, producing first-order, mixed first and zero-order, or zero-order kinetics as opposed to zero-order Michaelis–Menten kinetics (80). The rate of the reaction, \(v\), is proportional to the substrate concentration at the interface where

\[
v = \frac{V_{\text{max}}[S]}{K_m + [S]}
\]

The kinetics of immobilized lipases are also affected by the type of reactor used, since reactors differ in the amount of immobilized lipase used and in the method of substrate delivery, product removal, and degree of mixing.

V. ENZYMATIC INTERESTERIFICATION REACTORS

Reactors designed for immobilized enzyme reactions differ from one another based on several criteria. Reactors can be batch or flow-through systems and can differ in the degree of mixing involved during the reaction. For all reactor systems, the productivity of the system is defined as the volumetric activity \(\times\) the operational stability of the immobilized enzyme, with units of kilograms of product per liter of reactor.
volume per year. The volumetric activity is determined as the mass of product obtained per liter of reactor per hour, whereas the operational stability is the half-life of the immobilized enzyme (80). The most common reactor systems used include fixed bed, batch, continuous stirred tank, and membrane reactors.

A. Fixed Bed Reactor

A fixed bed reactor is a form of continuous flow reactor, where the immobilized enzyme is packed in a column or as a flat bed, and the substrate and product streams are pumped in and out of the reactor at the same rate. The main advantages of fixed bed reactors are their easy application to large scale production, high efficiency, low cost, and ease of operation. A fixed bed reactor also provides more surface area per unit volume than a membrane reactor system (7). A model fixed bed reactor for interesterification would consist of two columns in series: one for the reaction and a precolumn for fat-conditioning steps such as incorporation of water. Reservoirs attached to the columns would contain the feed streams and product streams. A pump would be required to keep the flow rate through the system constant, and the system would have to be water-jacketed to keep the reaction temperature constant (Fig. 15).

Since water is required in minimal amounts for hydration of the enzyme during the reaction, the oil is first passed through a precolumn containing water-saturated silica or molecular sieves, which would allow the oil to become saturated with sufficient water to allow progression of the interesterification reaction without increasing the rate of hydrolysis. Interesterification in a fixed bed reactor can lead to increases in product formation through increased residence time in the reactor. Complete conversion to products will never be achieved, and with an increase in product levels, a loss in productivity will occur (89). Using a fixed bed reactor with a silica pre-

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**Figure 15** Fixed bed reactor for immobilized lipase–catalyzed interesterification.
column for water saturation of the oil phase, Posorske et al. (89) produced a cocoa butter substitute from palm stearin and coconut oil. These authors found that decreasing the flow rates to increase the total product concentration caused a decrease in productivity. Decreasing the flow rates to increase product levels from 20% to 29% leads to a significant decrease in productivity. Fixed bed reactors are more efficient than batch reactors but are prone to fouling and compression. Dissolution of the oil in an organic solvent to reduce viscosity for flow through the packed bed may be required (89). In addition, the substrate has to be treated to remove any particulates, inhibitors, and poisons that can build up over time and inactivate the lipase (8). Macrae (9) found that after treatment of palm oil midfraction and stearic acid to remove particulates, inhibitors, and poisons, acidolysis reached completion after 400 hours and there was not appreciable loss in lipase activity even after 600 hours of operation. Wisdom et al. (91) performed a pilot scale reaction using a 2.9-L fixed bed reactor to esterify shea oleine with stearic acid. It was found that with high-quality substrates, only a small loss of activity was exhibited after 3 days with the production of 50 kg of product. However, when a lower grade shea oil was used, there was rapid inactivation of the lipase.

The kinetics of a packed bed reactor are assumed to be the same as for a soluble lipase, where

\[
\frac{dS}{dt} = \frac{V_{\text{max}}[S]}{K_m + [S]}
\]

This can be rearranged and integrated to

\[
[S_0]X = K_m' \ln(1 - X) + \frac{k_{\text{cat}}E_T}{Q}
\]

where \([S_0]\) is the initial substrate concentration, \(X\) is the fraction of substrate that has been converted to product at any given time \((1 - [S]/[S_0])\), \(Q\) is the volumetric flow rate, and \(E_T\) is the total number of moles of enzyme present in the packed bed (80,92). The residence time, \(\tau\), is based on the porosity of the packed bed and is defined as (93):

\[
\tau = \frac{V_{\text{tot}}}{P Q}
\]

where \(V_{\text{tot}}\) is the volume of the reactor, \(P\) is the porosity of the bed, and \(Q\) is the flow rate of the substrate.

The porosity of the bed in a fixed bed reactor can produce internal transfer limitations. Ison et al. (84) studied the effects of pore size on lipase activity in a fixed bed reactor using Spherosil with a mean pore size of 1480 Å and Duolite with a mean pore size of 190 Å. The larger pore size of the Spherosil was found to produce a decrease in lipase activity. This loss in activity was due to the higher degree of enzyme loading during immobilization, making some of the lipase inaccessible to substrate. With the smaller pore size of Duolite, the lipase was immobilized only on the surface of the support, eliminating internal mass transfer limitations.

### B. Stirred Batch Reactor

A stirred batch reactor is a common system used in laboratory experiments with lipase-catalyzed interesterification due to its simplicity and low cost. No addition
and removal of reactants and products is performed except at the initial and final stages of the reaction (Fig. 16). The equation to characterize the kinetics of a stirred batch reactor is:

\[
[S_0]X - K_m \ln(1 - X) = \frac{k_{cat}E_t t}{V}
\]

where \([S_0]\) is the initial substrate concentration, \(X\) is the fraction of substrate converted to product at any given time \((1 - [S]/[S_0])\), \(t\) is the reaction time, \(E_t\) is the total number of moles of enzyme present in the reactor, and \(V\) is the volume of the reactor.

Kurashige (94) found that a batch reactor was useful in reducing the diacylglycerol content in palm oil by converting existing diacylglycerols and free fatty acids to triacylglycerols. Using lipase coadsorbed with lecithin on Celite under vacuum to keep the water content below 150 ppm, the author was able to increase the triacylglycerol content from 85% to 95% in 6 hours. The rate of conversion in a stirred batch reactor decreases over time since there is a high initial level of substrate, which is reduced over time, with conversion to product. In order to maintain the same rate of conversion throughout the reaction, it would be necessary to add more immobilized enzyme to the reaction mixture (80). A stirred batch reactor has the advantage of being relatively easy to build and free enzymes can be used, but it has the disadvantage that, unless immobilized, the enzyme cannot be reused. As well, a larger system or longer reaction times are required to achieve equivalent degrees of conversion in comparison with other systems, and side reactions can be significant (63). Macrae (9) used a batch reactor to produce cocoa butter equivalents from the interesterification of palm oil midfraction and stearic acid. While product yields were high, by-products such as diacylglycerols and free fatty acids were formed. Therefore, it was necessary to isolate the desired triacylglycerols products using fat fractionation techniques.

C. Continuous Stirred Tank Reactor

A continuous stirred tank reactor combines components of both fixed bed and batch reactors. It is an agitated tank in which reactants and products are added and removed at the same rate, while providing continuous stirring to eliminate mass transfer lim-

![Stirred Batch Reactor](image)

**Figure 16** Stirred batch reactor for immobilized or free lipase–catalyzed interesterification.
itations encountered in fixed bed reactors (Fig. 17). Stirring also prevents the formation of temperature and concentration gradients between substrates or products. A continuous stirred tank reactor can be in the form of a tank with stirring from the top or bottom, or a column with stirring accomplished by propellers attached to the sides of the column (68). The kinetics for a continuous stirred tank reactor, developed by Lilly and Sharp (95), first encompass the substrate balance in the system as

\[ Q[S_i] - Q[S_0] = \frac{dS}{dt} V \]

where \( Q \) is the flow rate, \( [S_i] \) is the initial substrate concentration entering the reactor, \( [S_0] \) is the substrate concentration leaving the reactor, and \( V \) is the steady-state liquid volume in the tank.

Rearrangement gives:

\[ [S_0]X - K_{in}^\prime \left( \frac{X}{1-X} \right) = \frac{k_{cat}E_T}{Q} \]

where \([S_0]\) is the initial substrate concentration, \( X \) is the amount of substrate converted to product at any particular time \((1 - [S]/[S_0])\), \( Q \) is the flow rate, \( E_T \) is the total number of moles of enzyme present in the reactor.

The main disadvantages of continuous stirred tank reactors are the higher power costs associated with continuous stirring, the possibility of breaking up support particles with agitation, and the requirement for a screen or filter at the outlet to prevent losses of the immobilized lipase (7,80).

**D. Membrane Reactors**

Immobilization of enzymes onto semipermeable membranes is an attractive alternative for lipase-catalyzed interesterification reactions. Membrane reactors involve two-phase systems, where the interface of two phases is at a membrane. The advantages of membrane systems are reduced pressure drops, reduced fluid channeling, high effective diffusivity, high chemical stability, and a high membrane surface area to volume ratio (90). Membranes are commonly produced in the form of a bundle of hollow fibers and can be hydrophilic of hydrophobic in nature. Materials used in membrane systems are polypropylene, polyethylene, nylon, acrylic resin, and poly-

![Figure 17](image.png)  
**Figure 17** Continuous stirred tank reactor for immobilized lipase–catalyzed interesterification.
vinyl chloride. In a membrane such as microporous polypropylene, the pores have dimensions of 0.075 by 0.15 μm and the fibers have an internal diameter of 400 μm, providing 18 m² of surface area per gram of membrane (82). With a hydrophilic membrane such as cellulose, the oil phase circulates through the inner fiber side whereas the aqueous components circulate on the shell side (63). Immobilization of lipase can be accomplished by submerging the fibers in ethanol, rinsing them in buffer, then submerging them in lipase solution (82). Another method involves dispersing the enzyme in the oil phase and using ultrafiltration to deposit the lipase on the inner fiber side. One of the substrates can diffuse through the membrane toward the interface where the enzyme is immobilized. van der Padt et al. (63) used hollow fibers made from cellulose to perform glycerolysis of decanoic acid. Using a hydrophilic membrane bioreactor, the lipase activity was similar to the activity in emulsion systems. The hydrophilic membrane was found to be more effective for glycerolysis since the lipase was immobilized on the oil phase side, with the membrane preventing it from diffusing into the glycerol phase and being lost. Hoq et al. (96) used a hydrophobic polypropylene membrane to esterify oleic acid and glycerol. The lipase was adsorbed on the glycerol side, resulting in the loss of some enzyme in this phase. Therefore, use of a hydrophobic membrane would require the addition of more lipase to prevent losses in activity (7,64). Membrane reactors have been used in glycerolysis and acidolysis reactions and have an advantage over more conventional stirred tank reactors in that the reaction and separation of substrates and product can be accomplished in one system. Having the substrates and products separated during the reaction is especially useful during the esterification reaction where water is produced. Hoq et al. (96,97) found that during esterification of oleic acid and glycerol, the excess water produced could be removed by passing the oleic acid stream through molecular sieves, thereby preventing losses in productivity from hydrolysis.

E. Fluidized Bed Reactor

Fluidized bed reactors are reactors in which the immobilized enzyme and support are kept suspended by the upward flow of substrate or gas at high flow rates (80) (Fig. 18). The advantages of fluidized bed reactors are that channeling problems are eliminated, there is less change in pressure at high flow rates and less coalescence of emulsion droplets. Also, particulates do not have to removed from the oil and there are no concentration gradients (7). The main disadvantage of fluidized bed reactors is that small concentrations of enzyme can be used since a large void volume is required to keep the enzyme and support suspended. Mojovic et al. (98) used a gas lift reactor to produce a cocoa butter equivalent by interesterifying palm oil midfraction. These authors immobilized lipase encapsulated in lecithin reverse micelles in hexane; the reaction in the gas lift reactor was more efficient than in a stirred batch reactor. Equilibrium was reached 25% earlier and productivity was 2.8 times higher in the gas lift reactor.

VI. FACTORS AFFECTING LIPASE ACTIVITY DURING INTERESTERIFICATION

In considering all of the factors involved in enzymatic interesterification, all components of the system must be examined; namely pH, water content, temperature, substrate composition, product composition, and lipase content.
A. **pH**

Lipases are only catalytically active at certain pHs, depending on their origin and the ionization state of residues in their active sites. While lipases contain basic, neutral, and acidic residues, the residues in the catalytic site are only active in one particular ionization state. The pH optima for most lipases lies between 7 and 9, although lipases can be active over a wide range of acid and alkaline pHs, from about pH 4 to pH 10 (50,99). For example, the optimum pH for lipase from *Pseudomonas* species is around 8.5, whereas fungal lipases from *Aspergillus niger* and *Rhizopus delemar* are acidic lipases (100). The effect of immobilization on the pH optimum of lipases is dependent on the partitioning of protons between the bulk phase and the microenvironment around the support and the restriction of proton diffusion by the support. If the lipase is immobilized on a polyanion matrix, the concentration of protons in the immediate vicinity of the support will be higher than in the bulk phase, thereby reducing the pH around the enzyme in comparison with the pH of the bulk phase. Since there is a difference in the perceived pH of the solution as measured by the pH of the bulk phase, the lipase would exhibit a shift in pH optimum toward a more basic pH. For instance, for a free lipase that has a pH optimum of 8.0, when immobilized on a polyanionic matrix, with the bulk solution at pH 8.0, the pH in the immediate vicinity of the lipase might be only 7.0. Therefore, while the reaction pH is 8.0, the lipase is operating at pH 7.0, which is below its optimum. The pH of the bulk solution would have to be increased to pH 9.0 to get the pH around the lipase to its optimum of 8.0. This phenomenon is only seen in solutions with ionized support and low ionic strength systems (101). If protons are produced in the course of interesterification, the hydrogen ion concentration in the Nernst layer can be higher than in the bulk phase, thereby decreasing the pH in the vicinity of the lipase. Running an interesterification reaction with lipases at a pH well removed from the optimum can lead to rapid inactivation of the enzyme.

B. **Temperature**

In general, increasing the temperature increases the rate of interesterification, but very high temperatures can reduce the reaction rates due to irreversible denaturation.
of the enzyme. Animal and plant lipases are usually less thermostable than extracellular microbial lipases (99). In a solvent-free system, the temperature must be high enough to keep the substrate in the liquid state (84,102). Temperatures do not need to be as high in systems containing organic solvents since they easily solubilize hydrophobic substrates. However, for food industry applications, where organic solvents are avoided, the reaction temperatures are usually higher. Sometimes the temperature has to be increased to as high as 60°C to liquefy the substrate. Such high temperatures can seriously reduce the half-life of the lipase, although immobilization has been found to improve the stability of lipases under high temperature conditions. Immobilization fixes the enzyme in one conformation, which reduces the susceptibility of the enzyme to denaturation by heat. The optimal temperature for most immobilized lipases falls within the range of 30–62°C, whereas it tends to be slightly lower for free lipases (50). Immobilized lipases are more stable to thermal deactivation because immobilization restricts movement and can reduce the degree of unfolding and denaturation. Hansen and Eigtved (103) found that even at a temperature of 60°C, immobilized lipase from *Mucor miehei* has a half-life of 1600 hours.

C. Water Content and Water Activity

The activity of lipases at different water contents or water activity is dependent on the source of the enzyme. Lipases from molds seem to be more tolerant to low water activity than bacterial lipases. The optimal water content for interesterification by different lipases ranges from 0.04% to 11% (w/v), although most reactions require water contents of less than 1% for effective interesterification (15,50,104). The water content in a reaction system is the determining factor as to whether the reaction equilibrium will be toward hydrolysis or ester synthesis. Ester synthesis depends on low water activity. Too low a water activity prevents all reactions from occurring because lipases need a certain amount of water to remain hydrated, which is essential for enzymatic activity (34,105). As stated previously, lipases tend to retain the greatest degree of original activity when immobilized on hydrophobic supports. When the immobilized lipase is contacted with an oil-in-water emulsion, the oil phase tends to associate with and permeate the hydrophobic support, so that there is no aqueous shell surrounding the enzyme and support. It can be assumed that there is an ordered hydrophobic network of lipid molecules surrounding the support. Any water that reaches the enzyme for participation in hydrolysis and interesterification reactions must diffuse there from the bulk emulsion phase. Therefore, to avoid diffusional limitations, the oil phase must be well saturated with water (50). Too much water can inhibit interesterification, probably due to decreased access of hydrophobic substrates to the immobilized enzyme. Abraham et al. (106) found that in a solvent-free system, interesterification dominated hydrolysis up to a water-to-lipase ratio of 0.9, after which hydrolysis became the predominant reaction. During interesterification, the reaction equilibrium can be forced away from ester synthesis due to accumulation of water, 1 mol of which is produced for every mole of ester synthesized during the reaction. The equilibrium can be pushed back toward ester synthesis by continuous removal of water produced during the reaction. Water activity can be kept constant by having a reaction vessel with a saturated salt solution in contact with the reaction mixture via the gas phase in order to continuously remove the water produced in the course of interesterification. Another method of water activity
control that has proven useful with interesterification reactions is the use of silicone tubing containing the salt solution, immersed in the reaction vessel. Water vapor can be transferred out of the reaction system across the tubing wall and into the salt solution (107). A very simple method for water removal involves adding molecular sieves near the end of the reaction, or running the reaction under a vacuum so that the water produced is continuously removed, while still allowing the lipase to retain its water of hydration (44,94,108). Kurashige (94) ran an effective interesterification reaction with less than 150 ppm water maintained by running the reaction under vacuum.

D. Enzyme Purity and Presence of Other Proteins

During immobilization, adsorption of protein to surface-active supports is not limited solely to lipases. Other protein sources in the lipase solution can be adsorbed, and this can have an effect on the loading and activity of the immobilized enzyme. Use of a pure lipase solution for immobilization has been found to reduce activity of the lipase, whereas the presence of other proteins on the support can increase the activity of the immobilized lipase (91). Nonprotein sources of contamination during immobilization are usually not a problem because the lipase is preferentially adsorbed to the support.

E. Substrate Composition and Steric Hindrance

The composition of the substrate can have an effect on the rate of hydrolysis and interesterification by lipase. The presence of a hydroxyl group in the sn-2 position has a negative inductive effect, so that triacylglycerols are hydrolyzed at a faster rate than diacylglycerols, which are hydrolyzed at a faster rate than monoacylglycerols (11). While the nucleophilicity of substrate is important to the rate of reaction, steric hindrance can have a much greater negative effect. If the composition of the substrate is such that it impedes access of the substrate to the active site, any improvements in the nucleophilicity will not improve the activity (109).

The conformation of the substrate can also have an effect on the rate of reaction. The hydrophobic tunnel in the lipase accepts aliphatic chains and aromatic rings more easily than branched structures (11,44). For example, using carboxylic acids of differing chain lengths, Miller et al. (44) found that increasing the acyl group chain length up to seven carbons increased the esterification rate for lipase from *Mucor miehei*.

Oxidation of substrates, especially PUFAs, is possible and can cause inhibition and a decrease in activity of lipases, especially in reactions containing organic solvents. Inhibition is seen at hydroperoxide levels greater than 5 mequiv/kg oil and is attributed to the breakdown of hydroperoxides to free radicals (110). Therefore, before running interesterification reactions, especially in flow-through systems such as fixed bed reactors which are more susceptible to poisoning and inactivation, oils containing high levels of PUFAs must be highly refined (89).

F. Surface-Active Agents

The presence of surface-active agents used during the immobilization process can improve lipase activity during interesterification. The addition of lecithin or sugar
esters as surface active agents during the immobilization process can increase activity 10-fold when the preparation is used under microaqueous conditions (19). In contrast, using surface-active agents to form an emulsion can dramatically decrease the rate if interesterification because they prevent contact between the lipase and substrate (111). Adsorption at the interface can be inhibited by the presence of other nonsubstrate molecules, such as proteins. The presence of proteins other than lipase at the interface reduces the ability of the lipase to bind to the interface. Addition of protein in the presence of lipase can cause desorption of lipase from the interface.

Phospholipids, such as phosphatidylcholine, phosphatidylethanolamine, and phosphatidylinositol, can be found as minor components in oil, in quantities of 0.1–3.2%. The presence of phospholipids can have a negative effect on lipase activity. The initial rate of reaction can be decreased due to initial competition between phosphatidylcholine and the triacylglycerols for the active site of the lipase. Phosphatidylethanolamine seems to have the most inhibitory effect on lipase action possibly due to the presence of the amine group. Due to their effects, the phospholipid content of oils must be less than 500 ppm in order to prolong the half-life of immobilized lipases during interesterification (112).

G. Product Accumulation

During interesterification of two triacylglycerols, the production of monoacylglycerols and diacylglycerols can lead to an increase in the rate of reaction, whereas the presence of high levels of free fatty acids can inhibit the initial hydrolysis of triacylglycerols (51). In lipase-catalyzed interesterification, where hydrolysis is extensive, or in acidolysis reactions, the level of free fatty acids can have an impact on the rate of the reaction. During acidolysis of butter oil with undecanoic acid, Elliott and Parkin (65) reported that concentrations of undecanoic acid greater than 250 mM decreased the activity of porcine pancreatic lipase. Inhibition of lipase activity by free fatty acids agrees with the Michaelis–Menten model for uncompetitive inhibition by a substrate (65):

$$

\nu = \frac{V_{\text{max}}[S_0]}{[S_0] + \frac{[S_0]}{K_i} + K_m}

$$

where $S_0$ is the initial free fatty acid concentration, $K_i$ is the inhibition constant, and $K_m$ is the Michaelis constant.

The loss of activity by lipase in the presence of high concentrations of free fatty acids has been attributed to several factors. High levels of free fatty acids would produce high levels of free or ionized carboxylic acid groups, which would acidify the microaqueous phase surrounding the lipase or cause desorption of water from the interface. Also, with short and medium chain fatty acids, there could be partitioning of fatty acids away from the interface into the surrounding water shell due to their increased solubility in water. This would limit access by the substrate to the interface (113). Kuo and Parkin (113) found that there was less inhibition when longer chain fatty acids, such as C13:0 and C17:0, were used during acidolysis, compared with C5:0 and C9:0. The decrease in lipase activity was attributed to both increased solubility of the short chain fatty acids in and acidification of the aqueous phase.
VII. CONCLUSIONS

Despite the benefits of using lipase-catalyzed interesterification, it is unlikely that it will replace chemical interesterification in the future. This is due to the higher cost associated with enzymatic interesterification and the low cost of products, such as margarines and shortenings, that are currently being produced using chemical interesterification. The main attraction of lipase-catalyzed interesterification reactions is in the specificities of individual lipases and their application to the development of novel fats and oils that cannot be produced by chemical means. Future applications will involve continued development of reduced-calorie products, enriched lipids, and structured lipids. In addition, research will continue in the area of the characterization of fatty acid specificities of new lipases particularly in the identification of 2-specific lipases. In order for any of these new applications to be useful in the food industry, scale-up studies simulating industrial processes are necessary.

REFERENCES


Structured Lipids

CASIMIR C. AKOH

The University of Georgia, Athens, Georgia

I. INTRODUCTION

A. What Are Structured Lipids?

In a broad sense, structured lipids (SLs) are triacylglycerols that have been modified by incorporation of new fatty acids, restructured to change the positions of fatty acids, or the fatty acid profile, from the natural state, or synthesized to yield novel triacylglycerols (TAGs). This definition includes the topics covered in Chapters 23, 24, 25, and 30. The fatty acid profiles of conventional TAGs are genetically defined and unique to each plant or animal species. In this chapter, SLs are defined as TAGs containing mixtures of fatty acids (short chain and/or medium chain, plus long chain) esterified to the glycerol moiety, preferably in the same glycerol molecule. Figure 1 shows the general structure of SLs; their potency increases if each glycerol moiety contains both short and/or medium chain and long chain fatty acids. SLs combine the unique characteristics of component fatty acids such as melting behavior, digestion, absorption, and metabolism to enhance their use in foods, nutrition, and therapeutics. Individuals unable to metabolize certain dietary fats or with pancreatic insufficiency may benefit from the consumption of SL.

Structured lipids are often referred to as a new generation of fats that can be considered as “nutraceuticals”: food or parts of food that provide medical or health benefits, including the potential for the prevention and/or treatment of disease (1). Sometimes, they are referred to as functional foods or in the present context, as functional lipids. “Functional foods” is a term used to broadly describe foods that provide specific health benefits. Medical foods (medical lipids) are foods (lipids) developed for use under medical supervision to treat or manage particular disease or nutritional deficiency states. Other terms used to describe functional foods are phys-
B. Rationale for Structured Lipid Development

Over the past 39 years, long chain triacylglycerols (LCTs), predominantly soybean and safflower oils, have been the standard lipids used in making fat emulsions for total parenteral nutrition (TPN) and enteral administration. The emulsion provides energy and serves as a source of essential fatty acids (EFAs). However, long chain fatty acids (LCFAs) are metabolized slowly in the body. It was then proposed that medium chain triacylglycerols (MCTs) may be better than LCTs because the former are readily metabolized for quick energy. MCTs are not dependent on carnitine for transport into the mitochondria. They have higher plasma clearance, higher oxidation rate, improved nitrogen-sparing action, and less tendency to be deposited in the adipose tissue or to accumulate in the reticuloendothelial system (RES). One major disadvantage of using MCT emulsions is the lack of essential fatty acids (18:2n-6).

In addition, large doses of MCTs can lead to the accumulation of ketone bodies, a condition known as metabolic acidosis or ketonemia. It was suggested that combining MCTs and LCTs in the preparation of fat emulsions enables utilization of the benefits of both TAGs and may be theoretically better than pure LCT emulsions. An emulsion of MCTs and LCTs is called a physical mixture; however, a physical mixture is not equivalent to an SL. When MCTs and LCTs are chemically interesterified, the randomized product is called an SL. SLs are expected to be rapidly cleared and metabolized compared to LCTs.

For an SL to be beneficial, a minimum amount of LCFA is needed to meet essential fatty acid requirements. With the SL, LCFAs, medium chain fatty acids (MCFAs) and/or short chain fatty acids (SCFAs) can be delivered without the associated adverse effects of pure MCT emulsions. This is especially important when intravenous administration is considered (2,3). TAGs containing specific balances of medium chain, n-3, n-6, n-9, and saturated fatty acids can be synthesized to reduce serum low density lipoprotein (LDL) cholesterol and TAG levels, prevent thrombosis, improve immune function, lessen the incidence of cancer, and improve nitrogen balance (1,4). Although physical mixtures of TAGs have been administered to patients, an SL emulsion is more attractive because of the modified absorption rates of the SL molecule. Figure 2 shows the difference between a physical mixture of two triacylglycerols and SL pairs of molecular species.

SLs can be manipulated to improve their physical characteristics such as melting points. SLs are texturally important in the manufacture of plastic fats such as margarines, modified butters, and shortenings. Caprenin, a structured lipid produced.
by Procter & Gamble Company (Cincinnati, OH) consists of C8:0-C10:0-C22:0; it has the physical properties of cocoa butter but only about half the calories. Benefat™, originally produced as Salatrim (see Sec. II.B.2), consists of short chain (C2:0-C4:0) and long chain (C18:0) fatty acids. Both products can be used as cocoa butter substitutes. Currently, they are manufactured through a chemical transesterification process. Because of the low caloric value of the SCFAs and the partial absorption of stearic acid on Salatrim, this product has strong potential for use as a low-calorie fat substitute in the future. The caloric content of Caprenin and Benefat is about 5 kcal/g (vs. 9 kcal/g for a regular TAG). These SLs can also be manipulated for nutritive and therapeutic purposes, targeting specific diseases and metabolic conditions (4). In the construction of SLs for nutritive and therapeutic use, it is important that the function and metabolism of various fatty acids be considered. This chapter focuses mainly on SLs and MCTs, emphasizing the use of enzymes for SL synthesis as an alternative to chemical processing.

II. PRODUCTION OF STRUCTURED LIPIDS

A. Sources of Fatty Acids for Structured Lipid Synthesis

Structured lipids have been developed to optimize the benefit of fat substrate mixtures (5). A variety of fatty acids are used in the synthesis of SLs, taking advantage of the functions and properties of each to obtain maximum benefits from a given SL. These fatty acids include short chain fatty acids, medium chain fatty acids, polyunsaturated fatty acids, saturated long chain fatty acids, and monounsaturated fatty acids. Table 1 gives the suggested levels of some of these fatty acids in SLs intended for clinical applications. The component fatty acids and their position in the TAG molecule determine the functional and physical properties, the metabolic fate, and the health benefits of the SL. It is therefore appropriate to review the function and metabolism of the component fatty acids.
Table 1  Suggested Optimum Levels of Fatty Acids for Structured Lipids in Clinical Nutrition

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Levels and function</th>
</tr>
</thead>
<tbody>
<tr>
<td>n-3</td>
<td>2–5% to enhance immune function, reduce blood clotting, lower serum triacylglycerols, and reduce risk of coronary heart disease</td>
</tr>
<tr>
<td>n-6</td>
<td>3–4% to satisfy essential fatty acid requirement in the diet</td>
</tr>
<tr>
<td>n-9</td>
<td>monounsaturated fatty acid (18:1n-9) for the balance of long chain fatty acid</td>
</tr>
<tr>
<td>SCFA and MCFA*</td>
<td>30–65% for quick energy and rapid absorption, especially for immature neonates, hospitalized patients, and individuals with lipid malabsorption disorders</td>
</tr>
</tbody>
</table>

*Structured lipid containing short chain fatty acids (SCFAs) and/or medium chain fatty acids (MCFAs) as the main component.

Source Modified from Ref. 1.

1. Short Chain Fatty Acids

The SCFAs range from C2:0 to C6:0. They occur ubiquitously in the gastrointestinal tract of mammals, where they are the end products of microbial digestion of carbohydrates (6). In the human diet, SCFAs are usually taken in during consumption of bovine milk, which has a TAG mixture containing approximately 5–10% butyric acid and 3–5% caproic acid (7,8). Butyric acid is found in butterfat, where it is present at about 30% of the TAG (9). SCFAs, also known as volatile fatty acids, are more rapidly absorbed in the stomach than MCFAs because of their higher water solubility, smaller molecular size, and shorter chain length. Being hydrophilic, SCFAs have rates and mechanisms of absorption that are clearly distinguishable from those of lipophilic LCFAs (10). SCFAs are mainly esterified to the sn-3 position in the milk of cows, sheep, and goats (7). Under normal conditions, the end products of all carbohydrate digestion are the three major straight chain SCFAs: acetate, propionate, and butyrate (11,12); the longer SCFAs are generally found in smaller proportions except with diets containing high levels of sugar (13). Microbial proteolysis followed by deamination also produces SCFA.

Using synthetic TAGs, Jensen et al. (14) have shown that human pancreatic gastric lipase can preferentially hydrolyze sn-3 esters over sn-1 esters in the ratio of 2:1. This enzyme has also shown some hydrolytic specificity for short chain triacylglycerols (SCTs) and MCTs, although later studies (15) reported in vitro optimal conditions for the hydrolysis of LCFAs by gastric lipase. Pancreatic lipase has been reported to attack only the primary ester group of TAG, independent of the nature of fatty acid attached (16). Therefore, because of the positional and chain length specificity of the lipase, SCFAs attached to the sn-3 position of TAGs are likely to be completely hydrolyzed in the lumen of the stomach and small intestine. SCFAs are useful ingredients in the synthesis of low-calorie SLs such as Benefat because from heats of combustion, SCFAs are lower in caloric value than MCFAs and LCFAs. Examples of caloric values of SCFAs are as follows: acetic acid, 3.5 kcal; propionic acid, 5.0 kcal; butyric acid, 6.0 kcal; and caproic acid, 7.5 kcal.
2. Medium Chain Fatty Acids and Triacylglycerols

Medium chain triacylglycerols contain C6:0 to C12:0 fatty acids esterified to glycerol backbone. MCTs serve as an excellent source of medium chain fatty acids for SL synthesis. MCTs are used for making lipid emulsions either alone or by blending with LCTs for parenteral and enteral nutrition. The MCT structure is given in Figure 3. MCTs are liquid or solid at room temperature, and their melting points depend on the fatty acid composition. MCTs are used as carriers for colors, flavors, vitamins, and pharmaceuticals (17). MCFAs are commonly found in kernel oils or lauric fats; for example, coconut oil contains 10–15% C8:0 to C10:0 acid and is a raw material for MCT preparation (3). MCT is synthesized chemically by direct esterification of MCFA and glycerol at high temperature and pressure, followed by alkali washing, steam refining, molecular distillation, and further purification. Enzymatically, MCTs have been synthesized with immobilized *Mucor miehei* lipase in a solvent-free system (18). MCFAs have a viscosity of about 25–31 CP at 20°C and a bland odor and taste; as a result of the saturation of the fatty acids, they are extremely stable to oxidation (3). MCTs have a caloric value of 8.3 kcal compared to 9 kcal for LCTs. This characteristic has made MCTs attractive for use in low-calorie desserts. MCTs may be used in reduced-calorie foods such as salad dressings, baked goods, and frozen dinners (17).

MCTs have several health benefits when consumed in mixtures containing LCTs. Toxicological studies on dogs have shown that consuming 100% MCT emulsions leads to the development of adverse effects in dogs, which include shaking of the head and vomiting and defecation, progressing to a coma (19). It was theorized that these symptoms arose from elevated plasma concentration of MCFA or octanoate (19). Some advantages of MCFA/MCT consumption include the following: (a) MCFAs are more readily oxidized than LCFAs; (b) carnitine is not required for MCT transport into the mitochondria, thus making MCT an ideal substrate for infants and stressed adults (20); (c) MCFAs do not require chylomicron formation; and (d) MCFAs are transported back to the liver directly by the portal system. Absorption of SLs is discussed later in this chapter.

MCTs are not readily reesterified into TAGs and have better than twice the caloric density of proteins and carbohydrates, yet can be absorbed and metabolized as rapidly as glucose, whereas LCTs are metabolized more slowly (3). Feeding diets containing 20% and 30% lipid concentrations in weight maintenance studies indicate that MCTs may be useful in the control of obesity (21). MCTs appear to give satiety and satisfaction to some patients. Thermogenesis of MCT may be a factor in its very low tendency to deposit as depot fat (3).

Some reports suggest that MCTs can lower both serum cholesterol and tissue cholesterol in animals and man, even more significantly than conventional polyun-
saturated oils (22). However, a study by Cater et al. (23) showed that MCTs indeed raised plasma total cholesterol and TAG levels in mildly hypercholesterolemic men fed MCT, palm oil, or high oleic acid sunflower oil diets. A suggested mechanism for the cholesterol raising ability of MCTs is as follows: acetyl CoA, which is the end product of MCT oxidation, is resynthesized into LCFAs; the LCFAs then mix with the hepatic LCFA pool; and the newly synthesized LCFA may then behave like dietary LCFA. Also, the C8:0 may serve as precursor for de novo synthesis of LCFAs such as C14:0 and C16:0, which were detected in the plasma TAG (23). There were no differences in the high density lipoprotein (HDL) cholesterol concentrations among the subjects.

Evidence is pointing against the advisability of using MCTs in weight control because the level of MCTs (50%) required to achieve positive reduction is unlikely in human diet (24). An SL containing MCFA and linoleic acid bound in the TAG is more effective for cystic fibrosis patients than safflower oil, which has about twice as much linoleic acid as the SL (25). It appears that mobility, solubility, and ease of metabolism of MCFAs were responsible for the health benefits of the SL in these cases. In the SL, MCFAs provide not only a source of dense calories but also potentially fulfill a therapeutic purpose.

3. Omega-6 Fatty Acids

A common n-6 fatty acid is linoleic acid (18:2n-6). Linoleic acid is mainly found in most vegetable oils and in the seeds of most plants except coconut, cocoa, and palm nuts. Linoleic acids have a reducing effect on plasma cholesterol and an inhibitory effect on arterial thrombus formation (26). The n-6 fatty acids cannot be synthesized by humans and mammals and are therefore considered essential fatty acids (EFAs). The inability of some animals to produce 18:2n-6 is attributed to the lack of a Δ12 desaturase, required to introduce a second double bond in oleic acid. Linoleic acid can be desaturated further, and elongated to arachidonic acid (20:4n-6), which is a precursor for eicosanoid formation, as shown in Figure 4.

Essentiality of fatty acids was reported by Burr and Burr in 1929 (27). It is suggested that 1–2% intake of linoleic acid in the diet is sufficient to prevent biochemical and clinical deficiency in infants. Adults consume enough 18:2n-6 in the diet, and deficiency is not a problem. The absence of linoleic acid in the diet is characterized by scaly dermatitis, excessive water loss via the skin, impaired growth and reproduction, and poor wound healing (28). Nutritionists have suggested a 3–4% content of n-6 fatty acids in SLs to fulfill the essential fatty acid requirements of SLs (1).

4. Omega-3 Fatty Acids

Omega-3 fatty acids are also known as EFAs because humans, like all mammals, cannot synthesize them and therefore must obtain them from their diets. The n-3 fatty acids are represented by linolenic acid (18:3n-3), which is commonly found in soybean and linseed oils and in the chloroplast of green leafy plants. Other polyunsaturated n-3 fatty acids (n-3 polyunsaturated fatty acids, PUFAs) of interest in SL synthesis are eicosapentaenoic acid, 20:5n-3 (EPA) and docosahexaenoic acid, 22:6n-3 (DHA), which are commonly found in fish oils, particularly fatty fish. Children without enough n-3 PUFAs in their diet may suffer from neurological and visual
disturbances, dermatitis, and growth retardation (29). Therefore, n-3 PUFAs such as DHA must be included in their diet and in SL design.

Structured lipids containing n-3 PUFAs and MCFAs have been synthesized chemically by hydrolysis and random esterification of fish oil and MCTs. They have been shown to inhibit tumor growth and to improve nitrogen balance in Yoshida sarcoma-bearing rats (30). We have successfully used lipases as biocatalysts to synthesize position-specific SLs containing n-3 PUFAs with ability to improve immune function and reduce serum cholesterol concentrations (31,32). EPA is important in preventing heart attacks primarily because of its antithrombotic effect (33). It was also shown to increase bleeding time and to lower serum cholesterol concentrations (33). Studies with nonhuman primates and human newborns suggest that DHA is essential for the normal functioning of the retina and brain, particularly in premature infants (34). Other studies have shown that n-3 fatty acids can decrease the number and size of tumors and increase the time elapsed before the appearance of tumors (35).

The n-3 fatty acids are essential in growth and development throughout the life cycle of humans and therefore should be included in the diet. Nutritional experts consider a level of 2–5% of n-3 fatty acids optimum in enhancing immune function in SL as shown in Table 1. Polyunsaturated fatty acids of the n-3 series are antagonists of the arachidonic acid (20:4n-6) cascade (Fig. 5). The mode of action of fish oil n-3 PUFAs on functions mediated by n-6 PUFAs is summarized in Table 2 (36). The n-3 PUFAs inhibit tissue eicosanoid biosynthesis by preventing the action of δ-6 desaturase and cyclooxygenase/lipoxygenase enzymes responsible for the conversion of 18:2n-6 to 20:4n-6 and 20:4n-6 to eicosanoids, respectively. The amount of 18:2n-6 determines the 20:4n-6 content of tissue phospholipid pools and affects eicosanoid production. Eicosanoids are divided into prostanoids (prostaglandins, prostacyclins, and thromboxanes), which are synthesized via cyclooxygenase, and

---

**Figure 4** Pathway for eicosanoid biosynthesis.

\[\text{Linoleic Acid (18:2n-6)} \rightarrow \gamma-\text{Linolenic Acid (18:3n-3)} \rightarrow \text{Dihomo-\gamma-\text{Linolenic Acid (20:3n-6)}} \rightarrow \text{Leukotrienes} \rightarrow \text{Arachidonic Acid (20:4n-6)} \rightarrow \text{Tissue Phospholipids} \rightarrow \text{Prostaglandins} \rightarrow \text{Prostacyclin} \rightarrow \text{Endoperoxides} \rightarrow \text{Thromboxane} \]

---

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Figure 5  Pathways leading to the metabolism of dietary n-6 and n-3 polyunsaturated fatty acids.

Table 2  Mode of Action of n-3 PUFAs on Functions Mediated by n-6 PUFAs

| Impair uptake of n-6 polyunsaturated fatty acid (PUFA) |
| Inhibit desaturases, especially δ-6-desaturase |
| Compete with n-6 PUFAs for acyltransferases |
| Displace arachidonic acid (20:4n-6) from specific phospholipid pools |
| Dilute pools of free 20:4n-6 |
| Competitively inhibit cyclooxygenase and lipoxygenase |
| Form eicosanoid analogs with less activity or competitively bind to eicosanoid sites |
| Alter membrane properties and associated enzyme and receptor functions |

Source  Adapted from Ref. 36.
leukotrienes (hydroxy fatty acids and lipoxins), which are synthesized via lipooxygenase, as illustrated in Figure 4.

A proper balance of \( n-3 \) and \( n-6 \) fatty acids should be maintained in the diet and SL products. High concentrations of dietary 18:2\( n-6 \) may lead to increased production of immunosuppressive eicosanoids of the 2- and 4-series [prostaglandin E\(_2\) (PGE\(_2\)), thromboxane A\(_2\) (TXA\(_2\)), leukotriene B\(_4\) (LTB\(_4\))]. However, diets high in 20:5\( n-3 \) will inhibit eicosanoid production and reduce inflammation by increasing production of TXA\(_3\), prostacyclin (PGI\(_3\)), and LTB\(_5\). Diets including \( n-3 \) PUFAs also increase HDL-chol and interleukin-2 (IL-2) levels. On the other hand, they inhibit or decrease the levels of IL-1, LDL-chol, and very low density lipoprotein cholesterol (VLDL-chol).

5. Omega-9 Fatty Acids

The \( n-9 \) fatty acids or monounsaturates are found in vegetable oils such as canola, olive, peanut, and high-oleic sunflower as oleic acid (18:1\( n-9 \)). Oleic acid can be synthesized by the human body and is not considered an essential fatty acid. However, it plays a moderate role in reducing plasma cholesterol in the body (26). Oleic acid is useful in structured lipids for fulfilling the long chain triacylglycerol requirements of SLs as given in Table 1.

6. Long Chain Saturated Fatty Acids

Generally, saturated fatty acids are believed to increase plasma and serum cholesterol levels, but it has been claimed that fatty acids with chains 4–10 carbon atoms long do not raise cholesterol levels (37,38). Stearic acid has also been reported not to raise plasma cholesterol levels (39). TAGs containing high amounts of long chain saturated fatty acids, particularly stearic acid (18:0), are poorly absorbed in man partly because 18:0 has a melting point higher than body temperature; they exhibit poor emulsion formation and poor micellar solubilization (40). The poor absorption of saturated long chain triacylglycerols (40) makes them potential substrates for low-calorie SL synthesis. Indeed, Nabisco Foods Group used this property of stearic acid to make the group of low-calorie SLs called Salatrim (now Benefat) (see Sec. II.B.2), which consist of short chain aliphatic fatty acids and long chain saturated fatty acids, predominantly C18:0 (41). Caprenin, an SL produced by Procter & Gamble, contains C22:0, which is also poorly absorbed. An SL containing two behenic acids and one oleic acid has been used in the food industry to prevent chocolate bloom and to enhance fine crystal formation of palm oil and lard products (42).

B. Synthesis of Structured Lipids

1. Chemical Synthesis

Chemical synthesis of SLs usually involves hydrolysis of a mixture of MCTs and LCTs and then reesterification after random mixing of the MCFAs and LCFAs has occurred, by a process called transesterification (ester interchange). The reaction is catalyzed by alkali metals or alkali metal alkylates. This process requires high temperature and anhydrous conditions. Chemical transesterification results in desired randomized TAG molecular species, known as SLs, and in a number of unwanted products, which can be difficult to remove. The SL product consists of one (MLL,
Figure 6 Structure of Caprenin (caprocaprylobehenin) with three randomized acyl groups: $R_1$, $R_2$, $R_3$ = acyl part of capric acid, C10:0, caprylic acid, C8:0, and behenic acid, C22:0 in no particular order.
catalyzed interesterification of highly hydrogenated vegetable oils with TAGs of acetic and/or propionic and/or butyric acids (44). The product contains randomly distributed fatty acids attached to the glycerol molecule. Because of the random distribution of fatty acids, each preparation contains many molecular species. The ratio of SCFAs such as acetic, propionic, and butyric acids to LCFAs such as stearic acid can be varied to obtain SLs with physical and functional properties resembling those of conventional fats such as cocoa butter. The FDA accepted for filing in 1994 a GRAS affirmation petition by Nabisco.

Benefat is a low-calorie fat like Caprenin, with a caloric availability of 5 kcal/g. The caloric availability of C2:0, C3:0, C4:0, glycerol, and LCFA in the Benefat molecule are 3.5, 5.0, 6.0, 4.3, and 9.5 kcal/g, respectively. Stearic acid is poorly or only 50% absorbed (45), especially if it is esterified to the sn-1 and sn-3 positions of the glycerol. Acetyl and propionyl groups in Benefat are easily hydrolyzed by lipases in the stomach and upper intestine and readily converted to carbon dioxide (46). Benefat is intended for use in baking chips, chocolate-flavored coatings, baked and dairy products, dressings, dips, and sauces, or as a cocoa butter substitute in foods. The consistency of Benefat varies from liquid to semisolid, depending on the fatty acid composition and the number of SCFAs attached to the glycerol molecule. The structure of Benefat is given in Figure 7.

c. Others. Other commercially available chemically synthesized SLs and lipid emulsions are listed in Table 3. These include Captex, Neobee, and Intralipid (20%). Typical fatty acid profiles of selected SL products and MCTs are given in Table 4. Applications of these products will vary depending on the need of the patient or the function of the intended food product. Enzymes can be used to custom-produce SLs

Table 3 Commercial Sources of Structured Lipids and Lipid Emulsions

<table>
<thead>
<tr>
<th>Product</th>
<th>Composition</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caprenin</td>
<td>C8:0, C10:0, C22:0</td>
<td>Procter &amp; Gamble Co., Cincinnati, OH</td>
</tr>
<tr>
<td>Benefat</td>
<td>C2:0-C4:0, C18:0</td>
<td>Cultor Food Science, New York</td>
</tr>
<tr>
<td>Captex</td>
<td>C8:0, C10:0, C18:2</td>
<td>ABITEC Corp., Columbus, OH</td>
</tr>
<tr>
<td>Neobee</td>
<td>C8:0, C10:0, LCFA</td>
<td>Stepan Company, Maywood, NJ</td>
</tr>
<tr>
<td>Intralipid</td>
<td>20% soybean oil emulsion</td>
<td>KabiVitrum, Berkeley, CA</td>
</tr>
<tr>
<td>FE 73403</td>
<td>Fat emulsion of C8:0, C10:0, LCFA</td>
<td>Pharmacia AB, Stockholm, Sweden</td>
</tr>
</tbody>
</table>

LCFA, long chain fatty acid (may vary from C16:0 to C18:3n-3); FE, fat emulsion.
for specific applications. Unfortunately, many enzymatically synthesized SLs are not commercially available, although the potential is there. This technology needs to be commercialized.

3. Enzymatic Synthesis

a. Lipases in Fats and Oils Industry. Triacylglycerol lipases, also known as triacylglycerol acylhydrolases (EC 3.1.1.3), are enzymes that hydrolyze TAGs to DAGs, MAGs, free fatty acids (FFAs), and glycerol. They can catalyze the hydrolysis of TAGs and the transesterification of TAGs with fatty acids (acidolysis) or direct esterification of FFAs with glycerol (47–49). Annual sales of lipases presently account for only $20 million, which corresponds to less than 4% of the worldwide enzyme market estimated at $600 million (50). Two main reasons for the apparent misconception of the economic significance of lipases are as follows: (a) lipases have been investigated extensively as a route to novel biotransformation, and (b) the diversity of the current and proposed industrial applications of lipases by far exceeds that of other enzymes such as proteases or carbohydrases (51).

Although enzymes have been used for several years to modify the structure and composition of foods, they have only recently become available for large-scale use in industry, mainly because of the high cost of enzymes. However, according to enzyme manufacturers, progress in genetics and in process technology may now enable the enzyme industry to offer products with improved properties and at reduced costs (51). For lipases to be economically useful in industry, enzyme immobilization is necessary to enable enzyme reuse and to facilitate continuous processes. Immobilization of enzymes can simply be accomplished by mixing an aqueous solution of the enzyme with a suitable support material and removing the water at reduced pressure, after which small amounts of water are added to activate the enzyme. Suitable support materials for enzyme immobilization include glass beads, Duolite, acrylic resin, and Celite.

In spite of the obvious advantages of biological catalysis, the current level of commercial exploitation in the oleochemical industry is disappointing, probably be-

Table 4 Fatty Acid Composition of Typical Lipid Emulsions and Medium Chain Triacylglycerol

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>FE emulsion 73403</th>
<th>Intralipid 20% MCT</th>
</tr>
</thead>
<tbody>
<tr>
<td>8:0</td>
<td>27</td>
<td>—</td>
</tr>
<tr>
<td>10:0</td>
<td>10</td>
<td>—</td>
</tr>
<tr>
<td>12:0</td>
<td>—</td>
<td>1–2</td>
</tr>
<tr>
<td>16:0</td>
<td>7</td>
<td>13</td>
</tr>
<tr>
<td>18:0</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>18:1n-9</td>
<td>13</td>
<td>22</td>
</tr>
<tr>
<td>18:2n-6</td>
<td>33</td>
<td>52</td>
</tr>
<tr>
<td>18:3n-3</td>
<td>5</td>
<td>8</td>
</tr>
<tr>
<td>Other</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>

MCT, Medium chain triacylglycerol.
cause of the huge capital investments involved, and until recently, the high cost of lipase (51). The introduction of cheap and thermostable enzymes should tip the economic balance in favor of lipase use for the commercial production of SL and lipid modifications.

b. Mode of Action of Lipases. Triacylglycerol lipases are probably among the most frequently used enzymes in organic synthesis. This is in part because they do not require coenzymes and because they are stable enough in organic solvents at relatively high temperatures (52). Lipases act at the oil–water interface of heterogeneous reaction systems. This property makes them well suited for reactions in hydrophobic media. Lipases differ from esterases in their involvement of a lipid–water interface in the catalytic process (53). Some regions of the molecular structure responsible for the catalytic action of lipase are presumed to be different from those of ordinary enzymes that act on water-soluble substrates in a homogeneous medium (54). Because lipases work at substrate–water interfaces, a large area of interface between the water-immiscible reaction phase and the aqueous phase that contains the catalyst is necessary to obtain reasonable rates of interesterification (55). This is exemplified by the greater tendency for lipase to form off-flavors in homogenized milk than in unhomogenized milk.

Theoretical interpretations of the activation of lipase by interfaces can be divided into two groups: those assuming that the substrates can be activated by the presence of an oil–water interface, and those assuming that the lipase undergoes a change to an activated form upon contact with an oil–water interface. The first interpretation assumes higher concentrations of the substrate near the interface rather than in the bulk of the oil; and the second involves the existence of separate adsorption and catalytic sites for the lipase such that the lipase becomes catalytically active only after binding to the interface. More information on the action of microbial lipases is available in Chapter 26 of this book.

c. Enzymes in Organic Solvents. It is now commonly accepted that enzymes can function efficiently in anhydrous organic solvents. When enzymes are placed in an organic environment, they exhibit novel characteristics, such as altered chemo- and stereoselectivity, enhanced stability, and increased rigidity (56). Lipases have also been shown to catalyze peptide synthesis, since they can catalyze the formation of amide links while lacking the ability to hydrolyze them (57). Lipase can be used in several ways in the modification of triacylglycerols (48). In an aqueous medium, hydrolysis is the dominant reaction, but in organic media esterification and interesterification reactions are predominant. Lipases from different sources display hydrolytic positional specificity and some fatty acid specificity. The positional specificity is retained when lipases are used in organic media.

One application of lipases in organic solvents was their use as catalysts in the regio-specific interesterification of fats and oils for the production of TAGs with desired physical properties (58). Lipases can also be used in the resolution of racemic alcohols and carboxylic acids by the asymmetric hydrolysis of the corresponding esters. An example of stereoselectivity of lipases is the esterification of menthol by Candida cylindracea. This enzyme was shown to esterify L-menthol while being catalytically inactive with the D-isomer (59,60). Table 5 lists advantages of employing lipases in organic for the modification of lipids as opposed to aqueous media (61).
Table 5  Advantages of Lipase Modification of Lipids in Organic Solvents

<table>
<thead>
<tr>
<th>Advantage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increased solubility of nonpolar lipid substrates in organic solvents such as hexane and isooctane.</td>
</tr>
<tr>
<td>Shift of thermodynamic equilibria to the right in favor of synthesis over hydrolysis.</td>
</tr>
<tr>
<td>Reduction in water-dependent side reactions, since very little water is required by lipases in synthetic reactions.</td>
</tr>
<tr>
<td>Enzyme recovery is made possible by simple filtration of the powdered or immobilized lipase.</td>
</tr>
<tr>
<td>If immobilization is desired, adsorption onto nonporous surfaces (e.g., glass beads) is satisfactory; enzymes are unable to desorb from these surfaces in nonaqueous media.</td>
</tr>
<tr>
<td>Ease of recovery of products from low boiling point solvents.</td>
</tr>
<tr>
<td>Enhanced thermal stability of enzymes in organic solvents.</td>
</tr>
<tr>
<td>Elimination of microbial contamination.</td>
</tr>
<tr>
<td>Potential of enzymes to be used directly within a chemical process.</td>
</tr>
<tr>
<td>Immobilized enzyme can be reused several times.</td>
</tr>
</tbody>
</table>

Source Modified from Ref. 61.

d. Strategies for the Enzymatic Production of Structured Lipids. Various methods can be used for lipase-catalyzed production of SLs (4). The method of choice depends to a large extent on the type of substrates available and the products desired.

**Direct esterification.** Direct esterification can be used for the preparation of SLs by reacting free fatty acids with glycerol. The major problem is that the water molecules formed as a result of the esterification reaction must be removed as they are formed to prevent them from hydrolyzing back the product, leading to low product yield. Direct esterification, rarely used in SL synthesis, is presented in equation form as follows:

\[
\text{Glycerol} + \text{MCFA} + \text{LCFA} \xrightarrow{\text{lipase}} \text{SL} + \text{water}
\]

where MCFA = medium chain fatty acid, LCFA = long chain fatty acid, and SL = structured lipid moieties.

**Transesterification–acidolysis.** Acidolysis is a type of transesterification reaction involving the exchange of acyl groups or radicals between an ester and a free acid:

\[
\text{MCT} + \text{LCFA} \xrightarrow{\text{lipase}} \text{SL} + \text{MCFA}
\]

\[
\text{LCT} + \text{MCFA} \xrightarrow{\text{lipase}} \text{SL} + \text{LCFA}
\]

where MCT = medium chain triacylglycerol and LCT = long chain triacylglycerol.

**Figure 8** shows an example of acidolysis reaction (62), in this case between caprylic acid and triolein. Shimada et al. (63) used acidolysis reaction catalyzed by immobilized *Rhizopus delemar* lipase to synthesize an SL containing 22:6n-3 (DHA) and caprylic acids. Product isolation is easy after acidolysis. Free fatty acids are removed by distillation or by other appropriate techniques.

**Transesterification–ester interchange.** This reaction involves the exchange of acyl groups between one ester and another ester:
Figure 8  Reaction scheme showing acidolysis reaction in the synthesis of structured lipids from caprylic acid and triolein (62).

\[
\text{MCT} + \text{LCT} \xrightarrow{\text{lipase}} \text{SL} \\
\text{LCT} + \text{MCFAEE} \xrightarrow{\text{lipase}} \text{SL} + \text{LCFAEE} \\
\text{MCT} + \text{LCFAEE} \xrightarrow{\text{lipase}} \text{SL} + \text{MCFAEE}
\]

where MCFAEE = medium chain fatty acid ethyl ester and LCFAEE = long chain fatty acid ethyl ester. This method is widely used in lipid modifications and in the synthesis of SLs (4,47,64,65).

In a transesterification reaction, generally, hydrolysis precedes esterification. In all the preceding examples, short chain triacylglycerols (SCTs) or short chain fatty acids (SCFAs) can replace MCTs and MCFAs, respectively, or can be used in combination. Figures 9 and 10 give examples of the suggested strategies involving interchange reactions between a TAG (trilinolein) and a TAG (tricaprin) ester and between EPA ethyl ester and tricaprin, respectively. We have successfully used enzymes to synthesize position-specific SLs containing n-3 PUFAs with ability to improve immune function and reduce serum cholesterol (31,32).

e. Factors That Affect Enzymatic Process and Product Yield

Water. It is generally accepted that water is essential for enzymatic catalysis. This status is attributed to the role water plays in all noncovalent interactions. Water is responsible for maintaining the active conformation of proteins, facilitating reagent diffusion, and maintaining enzyme dynamics (66). Zaks and Klibanov (67) reported that for enzymes and solvents, tested enzymatic activity greatly increased with an increase in the water content of the solvent. The absolute amount of water required for catalysis for different enzymes varies significantly from one solvent to another (56). Hydration levels corresponding to one monolayer of water can yield active enzymes (68). Although many enzymes are active in a variety of organic solvents, the best nonaqueous reaction media for enzymatic reactions are hydrophobic, water-immiscible solvents (67,69,70). Enzymes in these solvents tend to keep the layer of essential water, which allows them to maintain their native configuration, and therefore catalytic activity.
Solvent type. The type of organic solvent employed can dramatically affect the reaction kinetics and catalytic efficiency of an enzyme. Therefore, the choice of solvent to be used in biocatalysis is critical. Two factors affecting this choice are the extent to which the solvent affects the activity or stability of the enzyme, and the effect of the solvent on the equilibrium position of the desired reaction (71). The equilibrium position in an organic phase is usually different from that in water because of differential solution of the reactants. For example, hydrolytic equilibrium is usually shifted in favor of the synthetic product because the product is less polar than the starting materials (71). The nature of the solvent can also cause inhibition or inactivation of enzymes by directly interacting with the enzyme. Here the solvent

Figure 9  Ester interchange reaction between two triacylglycerols, trilinolein and tricaprin, in the enzymatic production of structured lipids.

Figure 10  Ester interchange reaction in the production of structured lipids containing eicosapentaenoic acid (EPA) with tricaprin and EPA ethyl ester as substrates. An immobilized Candida antarctica lipase, SP 435, was the biocatalyst. Note EPA esterified to the sn-2 position.
alters the native conformation of the protein by disrupting hydrogen bonding and hydrophobic interactions, thereby leading to reduced activity and stability (72).

Lipases differ in their sensitivity to solvent type. An important solvent characteristic that determines the effect of solvent in enzymatic catalysis is the polarity of the solvent. Solvent polarity is measured by means of the partition coefficient of a solvent between octanol and water (73), and this is taken as a quantitative measure of polarity, otherwise known as log \( P \) value (74). The catalytic activity of enzymes in solvents with log \( P < 2 \) is usually lower than that of enzymes in solvents with log \( P > 2 \). This is because hydrophilic or polar solvents can penetrate into the hydrophilic core of the protein and alter the functional structure (75). They also strip off the essential water of the enzyme (67). Hydrophobic solvents are less able to remove or distort the enzyme-associated water and are less likely to cause inactivation of enzymes (61).

In choosing a solvent for a particular reaction, two important factors must be taken into consideration: the solubility of the reactants in the chosen solvent and the need for the chosen solvent to be inert to the reaction (61). Other factors that must be taken into account in determining the most appropriate solvent for a given reaction include solvent density, viscosity, surface tension, toxicity, flammability, waste disposal, and cost (61). A report by Akoh and Huang (62) on the effect of solvent polarity on the synthesis of SLs using IM 60 lipase from \textit{Rhizomucor miehei} showed that nonpolar solvents such as isooctane and hexane produced 40 mol\% of disubstituted SL, while a more polar solvent such as acetone produced 1.4% of the same SL. Claon and Akoh (76) found that with SP 435 lipase from \textit{Candida antarctica}, a higher log \( P \) value does not necessarily sustain a higher enzyme activity. Some experimentation is therefore necessary in selecting solvents for enzymatic reactions.

\( pH \). Enzymatic reactions are strongly \( pH \) dependent in aqueous solutions. Studies on the effect of \( pH \) on enzyme activity in organic solvents show that enzymes “remember” the \( pH \) of the last aqueous solution to which they were exposed (65,70). That is, the optimum \( pH \) of the enzyme in an organic solvent coincides with the \( pH \) optimum of the last aqueous solution to which it was exposed. This phenomenon is called \( pH \) memory. A favorable \( pH \) range depends on the nature of the enzyme, the substrate concentration, the stability of the enzyme, the temperature, and the length of the reaction (77).

\textit{Thermostability}. Temperature changes can affect parameters such as enzyme stability, affinity of enzyme for substrate, and preponderance of competing reactions (78). Thermostability of enzymes is a major factor the industry considers prior to commercialization of any enzymatic process, mostly because of the potential for saving energy and minimizing thermal degradation. Thermostability of lipases varies considerably with enzyme origin: animal and plant lipases are usually less thermostable than microbial extracellular lipases (49).

Several processes that lead to the irreversible inactivation of enzymes involve water as a reactant (79). This characteristic of enzymes makes them more thermostable in water-restricted environments such as organic solvents. Enzymes are usually inactivated in aqueous media at high temperatures. Several studies have been reported on the effect of temperature on lipase activity (64,76,80). Zaks and Klibanov (80), who studied the effect of temperature on the activity of porcine pancreatic lipase, showed that in aqueous solution at 100\(^\circ\)C, the lipase is completely inactivated within seconds, whereas in dry tributyrin containing heptanol, the lipase had a shelf
life at 100°C of 12 hours. These investigators concluded that in organic solvents, porcine pancreatic lipase remains rigid and cannot undergo partial unfolding, which causes inactivation. The heat stability of a lipase also depends on whether a substrate is present. This is because substrates remove excess water from the immediate vicinity of the enzyme, thus restricting its overall conformational mobility (81).

Most lipases in nonimmobilized form are optimally active between 30 and 40°C (82). Immobilization confers additional stability to the lipase compared to nonimmobilized lipase. Excellent reviews on the immobilization procedures and bioreactors for lipase catalysis were published recently (48,83,84). The immobilization support must possess the following properties: high surface area to allow maximum contact with enzyme, high porosity to allow good flow properties, high physical strength, solvent resistance, high flow properties, and chemical and microbiological inertness (85,86).

Other Factors. Other factors that affect yield of products are substrate molar ratio; enzyme source, activity, and load; incubation time; specificity of the enzyme to substrate type and chain length; and regiospecificity.

f. Chemical Versus Enzymatic Synthesis. The most useful property of lipases is their regio- and stereospecificity, which result in products with better defined and more predictable chemical composition and structure than those obtained by chemical catalysis. Potential advantages of using enzymes over chemical procedures may be found in the specificity of enzymes and the mild reaction conditions under which enzymes operate (87). Enzymes form products that are more easily purified and produce less waste, and thus make it easier to meet environmental requirements (87). Chemical catalysts randomize fatty acids in triacylglycerol mixtures and do not lead to the formation of specialty products with desired physicochemical characteristics (51). The specificities of lipase have classically been divided into five major types: lipid class, positional, fatty acid, stereochemical, and combinations thereof (81). Enzymes have high turnover numbers and are well suited for the production of chiral compounds important to the pharmaceutical industry.

Transesterification using sn-1,3 specific lipase results in SL products with fatty acid at the sn-2 fatty acids remaining almost intact. This is significant from a nutritional point of view because the 2-MAGs produced by pancreatic lipase digestion are the main carriers of fatty acids through the intestinal wall (88). Fatty acids esterified at the sn-2 position are therefore more efficiently absorbed than those at the sn-1 and sn-3-positions. A TAG containing an essential fatty acid at the sn-2 position and short or medium chain fatty acid in the sn-1 and sn-3 positions has the advantage of efficiently providing an EFA and a quick energy source (89).

Some studies have shown that the rate of autoxidation and melting properties of TAGs can be affected by the position of unsaturated fatty acids on the glycerol molecule (90,91). TAGs having unsaturated fatty acids at the 2-position of glycerol are more stable toward oxidation than those linked at the 1- and 3-positions.

The energy saved and minimizations of thermal degradation are probably among the greatest attractions in replacing the current chemical technology with enzyme biotechnology (51). Table 6 shows some of the potential advantages of the enzymatic approach to structured lipid design. Potential food applications of SL are listed in Table 7.
Table 6  Advantages of Enzymatic Approach to Structured Lipid Design

Position-specific SL (i.e., desirable fatty acids can be incorporated at specific positions of triacylglycerol).
Enzymes exhibit regioselectivity (discriminate based on bond to be cleaved), enantioselectivity (optical activity), chemoselectivity (based on functional group), and fatty acid chain length specificity.
Can design SL on case-by-case basis to target specific food or therapeutic use—custom synthesis.
Products with defined structure can be produced.
Novel products not possible by conventional plant breeding and genetic engineering can be obtained (e.g., by inserting specific fatty acid at the sn-2 position of glycerol molecule).
Mild reaction conditions.
Few or no unwanted side reactions or products.
Can control the overall process.
Ease of product recovery.
Add value to fats and oils.
Improve functionality and properties of fats.

4. Analysis of Structured Lipids

Figure 11 presents a purification and analysis scheme for enzymatically produced SLs. Method of analysis depends on whether the SL is synthesized by acidolysis or by interesterification reaction. The crude SL product can be analyzed with silica gel G or argentation AgNO₃ (based on unsaturation), thin-layer chromatography (TLC), gas-liquid chromatography (GLC) of the fatty acid methyl or ethyl esters for fatty acid profile, and by reversed phase high-performance liquid chromatography (RP-HPLC) of molecular species based on equivalent carbon number (ECN) or total carbon number (TCN). A typical HPLC chromatogram of SL products is shown in Figure 12.

Other methods of typical lipid analysis described in this book can be applied to studies of structured lipids. The choice of fractionation or purification technique depends on substrate or reactant types, products formed, overall cost, and whether a small-scale or large-scale synthesis was employed. The need for improved methodologies for the analysis of SCFA and MCFA components of SLs is emphasized here because of their volatility during extraction and GLC analysis.

Table 7  Potential Food Uses of Structured Lipids

Margarine, butter, spreads, shortening, dressings, dips, and sauces
Improve melting properties of fats
Cocoa butter substitute
Confectioneries
Soft candies
As reduced- or low-calorie fats (e.g., Caprenin, Benefat)
Baking chips, baked goods
Snack foods
Dairy products
Figure 11  Purification and analysis scheme for enzymatically produced structured lipids.

a. Stereospecific Analysis. Figure 13 shows the stereochemical configuration of a TAG molecule with sn notation indicating the stereochemical numbering system. The positional distribution of SFCA, MCFA, and LCFA on the glycerol moiety of SL is important in relation to the physical and functional properties of the SL, and its metabolism. As indicated below, the absorption and transport pathway of the SL depend somewhat on the fatty acid at the sn-2 position. In most vegetable oils, unsaturated fatty acids occupy the sn-2 position and saturated fatty acids are located

Figure 12  High-performance liquid chromatographic separation of structured lipid products from the reactants using a reversed phase column: SL1, structured lipid containing two medium chain fatty acids; SL2, structured lipid containing one medium chain fatty acid. Trilinolein and tricaprin were the reactants, and triolein was the internal standard.
Figure 13 Stereochemical configuration of triacylglycerols or structured lipids with $sn$ notation indicating stereochemical numbering of the carbon atoms of glycerol moiety. When the carbon in the 2-position is in the plane of the page and the 1- and 3-carbons behind the plane of the page, if the $-\text{OH}$ on the 2-position of glycerol is drawn to the left, the top carbon becomes 1 and the bottom becomes 3. Thus, a structured lipid with octanoic acid on the 1-position, and oleic acid on the 2-position and decanoic acid on the 3-position is named $sn$-glycerol-1-octanoate-2-oleate-3-decanoate.

in the $sn$-1 and $sn$-3 positions (92,93,95,96). The $sn$-2 position of TAG is determined by pancreatic lipase hydrolysis of the fatty acids at the $sn$-1 and $sn$-3 positions, followed by GLC analysis of the 2-MAG fatty acid methyl or ethyl ester. Detailed stereospecific analysis of the fatty acids at all three positions of the glycerol molecule was excellently reviewed by Small (93) and is not discussed in detail here. $^{13}$CNMR was used to determine acyl position of fatty acids on glycerol molecule (94).

Grignard reagent or Grignard degradation (97,98) is useful in obtaining the complete stereochemical structure of any TAG following pancreatic hydrolysis. In general, phospholipid derivatives (phosphatidylcholine, PC) of 1,2-DAG and 2,3-DAG are made by reacting with phospholipase A$_2$ (PLA$_2$). Since the $sn$-2 fatty acid is known, chemical analysis of the 2,3-diacyl-PC PLA$_2$ hydrolysis product gives the fatty acid at the $sn$-3 position. Similarly, chemical analysis of the 1,2-DAG hydrolysis product of PLA$_2$ gives the fatty acid at the $sn$-1 position. Alternatively, pancreatic hydrolysis of the 1,2-DAG followed by chemical analysis can give the fatty acid at the $sn$-1 position, since this enzyme is $sn$-1,3 specific.

III. ABSORPTION, TRANSPORT, AND METABOLISM OF STRUCTURED LIPIDS

The influence of TAG structure on lipid metabolism has been the subject of recent reviews and research efforts (92,93,99–101). SLs may be targeted for either portal or lymphatic transport. In one widely accepted pathway, C$_2$:0 to C$_{12}$:0 fatty acids are transported via the portal system and C$_{12}$:0 to C$_{24}$:0 via the lymphatic system (2). There is growing evidence that MCFAs may indeed be absorbed as 2-MAG, especially if they are esterified to the $sn$-2 position of the SL. The rate of hydrolysis at the $sn$-2 position of TAG is very slow, and as a result the fatty acid at this position remains intact as 2-MAG during digestion and absorption. Indeed, close to 75% of $sn$-2 position fatty acids are conserved throughout the process of digestion and absorption (102).

LCTs are partially hydrolyzed by pancreatic lipase and absorbed slowly as partial glycerides in mixed micelles (93). The resulting LCFAs are reesterified and incorporated into chylomicrons in the enterocyte, whereupon they enter the lymphatics to reach the general circulation through the thoracic duct. However, MCTs
are nearly completely hydrolyzed and absorbed faster, mainly as free fatty acids and rarely as 2-MAGs. These MCFAs are then transported as FFAs bound to serum albumin in portal venous blood.

Figure 14 shows a proposed modified pathway for MCT, LCT, and SL metabolism. The proposed metabolic routes for SL based on current evidence are indicated in italics. The metabolism of an SL is determined by the nature and position of the constituent fatty acids on the glycerol moiety. This may account for the differences in the pathway of absorption: lymphatics versus portal.

Evidence for lymphatic absorption of MCFAs and storage in adipose tissue is accumulating (103–107). Jensen et al. (107) observed the presence of more C10:0 than C8:0 fatty acids in the lymph of canine model fed an SL containing MCTs and fish oil versus its physical mixture, despite an overall ratio of C10:0 to C8:0 of 0.3 in their diets. Analysis of the SL molecular species revealed that MCFAs in lymph were present as mixed TAGs, suggesting that the MCFAs at the 2-position may account for the improved absorption. The 2-MAGs apparently were reesterified with endogenous or circulating LCFAs and subsequently absorbed through the lymphatic system. Also, feeding of high levels of MCTs can lead to lymphatic absorption and presence of MCFAs in the chylomicrons.

Enhanced absorption of 18:2n-6 was observed in cystic fibrosis patients fed SL containing LCFAs and MCFAs (89,108). Rapid hydrolysis and absorption of an SL containing MCFAs at the sn-1 and sn-3 positions and an LCFA at the sn-2 position have been reported (89,109,110). To improve the absorption of any fatty acid, its esterification to the 2-position of the glycerol moiety is suggested. Mok et al. (111) reported that the metabolism of an SL differs greatly from that of a similar physical mixture. The purported benefit of fish oil n-3 PUFAs can be attributed to their absorption as 2-MAGs. This factor is important in the construction of novel or designer SL molecules for food, therapeutic, and nutritional use.

Figure 14 Proposed modified metabolic pathways for medium chain and long chain triacylglycerols and structured lipids.
IV. NUTRITIONAL AND MEDICAL APPLICATIONS

Structured lipids can be synthesized to target specific metabolic effects or to improve physical characteristics of fats. An SL made from fish oil and MCTs was compared with conventional LCTs and found to decrease tumor protein synthesis, reduce tumor growth in Yoshida sarcoma-bearing rats, decrease body weight, and improve nitrogen maintenance (30). In addition, the effects of fish/MCT on tumor growth was synergistic with tumor necrosis factor (TNF). A similar study by Mendez et al. (112) compared the effects of a structured lipid (made from fish oil and MCFAs) with a physical mix of fish oil and MCTs and found that the SL resulted in improved nitrogen balance in animals, probably because of the modified absorption rates of SL. Gollaher et al. (113) reported that the protein-sparing action associated with SL administration are not seen when the structured lipids provide 50% of protein calories and suggested that the protein-sparing action of SLs may be dependent on the ratio of MCTs to LCTs used to synthesize the SL.

Jandacek et al. (89) demonstrated that a structured lipid containing octanoic acid at the 1- and 3-positions and a long chain fatty acid in the 2-position is more rapidly hydrolyzed and efficiently absorbed than a typical LCT. They proposed that the SL may be synthesized to provide the most desirable features of LCFAs and MCFAs for use as nutrients in cases of pancreatic insufficiency (89). Metabolic infusion of an SL emulsion in healthy humans showed that the capacity of these subjects to hydrolyze SL is at least as high as that to hydrolyze LCT (114). This finding is significant because of evidence of interaction and interference in the metabolism of LCT and MCT when both are present in a physical mix (115,116). An investigation into the in vivo fate of fat emulsions based on SL showed potential for use of SL as core material in fat emulsion–based drug delivery systems (117).

An SL made from safflower oil and MCFAs was fed to injured rats, and the animals receiving the structured lipid were found to have greater gain in body weight, greater positive nitrogen balance, and higher serum albumin concentration than controls receiving a physical mix (111). Enhanced absorption of 18:2n-6 was observed in cystic fibrosis patients fed structured lipids containing LCFAs and MCFAs (25). A mixed acid type of triacylglycerol composed of linoleic acid and MCFAs has been reported to improve immune functions (118), and evaluations in clinical nutrition are ongoing. However, a 3:1 admixture of MCT-LCT emulsions was reported to elevate plasma cholesterol concentrations compared to LCT emulsions in rats fed by intravenous infusion (119).

SL appears to preserve reticuloendothelial function while improving nitrogen balance as measured by the organ uptake of radiolabeled Pseudomonas in comparison to LCT (120). Long-term feeding studies with an SL containing MCFAs and fish oil fatty acids showed that SL modified plasma fatty acid composition, reflecting dietary intake and induced systemic metabolic changes that persisted after the diet was discontinued (121). An SL made by reacting tripalmitin with unsaturated fatty acids using an sn-1,3 specific lipase closely mimicked the fatty acid distribution of human milk was commercially developed for application in infant formulas under the trade name Betapol (122). HDL cholesterol decreased by 14% when a diet containing Caprenin as 40% of total calories was fed to healthy men, compared to no change in levels when an LCT diet was fed (43). Table 8 lists the potential and other reported benefits of SL (1,25,32,89,106,107,111,120,123–129).
Table 8  Potential and Reported Benefits of Structured Lipids

<table>
<thead>
<tr>
<th>Benefit</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Superior nitrogen retention</td>
<td>111</td>
</tr>
<tr>
<td>Preservation of reticuloendothelial system (RES) function</td>
<td>120</td>
</tr>
<tr>
<td>Attenuation of protein catabolism and the hypermetabolic stress response to thermal injury</td>
<td>123–125</td>
</tr>
<tr>
<td>Enhanced absorption of the fatty acid at the sn-2 position (e.g., 18:2n-6 cystic fibrosis patients)</td>
<td>25, 126</td>
</tr>
<tr>
<td>Reduction in serum TAG, LDL-cholesterol, and cholesterol</td>
<td>32, 106</td>
</tr>
<tr>
<td>Improved immune function</td>
<td>1, 127</td>
</tr>
<tr>
<td>Prevention of thrombosis</td>
<td>1</td>
</tr>
<tr>
<td>Lipid emulsion for enteral and parenteral feeding</td>
<td>127, 128</td>
</tr>
<tr>
<td>Calorie reduction</td>
<td>129</td>
</tr>
<tr>
<td>Improved absorption of other fats</td>
<td>89, 106, 107</td>
</tr>
</tbody>
</table>

Source  Modified from Ref. 4.

V. SAFETY AND REGULATORY ISSUES

The problem with consuming large doses of pure MCTs or their emulsions is the tendency to form ketone bodies (i.e., to induce metabolic acidosis). This outcome can be circumvented by using SLs or their emulsions. SL is safe and well tolerated in the body. Physiological and biochemical data suggest that SL emulsions, Intralipid 20%, and fat emulsion 73403 (Kabi Pharmacia AB, Stockholm, Sweden), when fed to postoperative patients, were rapidly cleared and metabolized (130). The safety of Benefat was assessed, and no significant clinical effects were reported in subjects consuming up to 30 g/day (131). Other studies also indicate that SLs are safe (132).

SLs that provide fewer calories (<9 kcal/g) than conventional TAGs (9 kcal/g) pose a great challenge to the FDA and other regulatory agencies around the world. These SLs include Caprenin, Benefat, and Captex. The issue is complicated by the labeling requirements for reduced fats. The big problem is how to establish uniform digestibility and absorbability coefficients for all available and soon-to-be-available SL molecules and other fat substitutes (133). The current dietary guidelines recognize total fat and saturated fat, but not digestibility coefficients. The FDA needs to develop new guidelines for SLs and genetically engineered vegetable oils or to modify existing guidelines for TAGs to reflect the new generation of fats. FDA accepted for filing the GRAS petition for Benefat/Salatrim in 1994.

VI. PERSPECTIVES

This chapter discussed the currently available structured lipids, methods of synthesis, raw materials considerations, and SL applications. An understanding of the functional properties and metabolic fate of the component fatty acids will aid in the synthesis of new SL molecules with beneficial end-use properties. The key to efficient absorption rests on the stereochemical structure of the SL. With this in mind, the outlook and potential for commercialization (Table 9) of the enzymatic process is bright.
Table 9  Factors That Affect Outlook for Commercialization of the Enzymatic Process

| Specialty needs or niche market as food ingredient, fine chemical use, nutritional supplement, enteral and parenteral feeding. |
| Cost of enzymatic versus chemical process and product yield. |
| Ease with which the enzymatic process can be scaled up. |
| Position specific with enzymes versus randomized products with chemical synthesis. |
| Consumer preference: natural versus synthetic products. |
| Cost-benefit assessment: investment and potential returns. |
| Catalyst reuse: immobilized enzyme can be reused several times without significant loss of activity. |
| Side or unwanted products of the reaction. |
| Processing costs to obtain products of high purity. |
| Regulation by the FDA (time-consuming process). |
| Competition with genetically engineered crops that produce structured lipids (e.g., high lauric acid canola oil). |

Enzymes allow scientists to design SLs intended for various applications, which may include treatment of cystic fibrosis patients; individuals with pancreatic insufficiency; acquired immune deficiency syndrome (AIDS) patients, who need to boost their immune system by consuming SLs containing 20:5n-3 at the sn-2 position; stressed and/or septic and hospital patients; and preterm infants. Potential nonmedical applications include foods and nutritional supports. SLs will continue to play a role in enteral and parenteral nutrition.

More research is needed on the effect of all lipid emulsions, especially SL emulsions, on the immune system. The notion that MCTs do not go via lymphatic transport is becoming less acceptable in the scientific community. The use of enzymes in constructing SLs destined for either portal or lymphatic transport will greatly enhance our knowledge on how SLs are metabolized. Chemical synthesis will lead to randomized SLs. Since the position and type of fatty acids in the TAG is key to their metabolism, the best alternative to chemical synthesis is the use of lipases. More applications of SLs in our regular diets is encouraged, meaning that food technologists need to explore this further. Genetic engineering of vegetable oil producing plants as covered in Chapter 30 will play a role in future commercial availability of SL.

REFERENCES


Biosynthesis of Fatty Acids and Storage Lipids in Oil-Bearing Seed and Fruit Tissues of Plants

KIRK L. PARKIN

University of Wisconsin–Madison, Madison, Wisconsin

I. INTRODUCTION

Upon arriving at this chapter, the reader may immediately react by questioning the relevance of fatty acid and triacylglycerol biosynthesis to mainstream food science. “Is that not better left to the biochemists?” might be a common retort. One challenge assumed by the author in agreeing to prepare this chapter is to convince the reader that it is important for food scientists to have a working knowledge of lipid biosynthesis. The next paragraph is intended to provide the initial fiber for this argument. The conclusion of this chapter (Sec. VI) strives to reinforce the argument by identifying examples of how food science research can be put to use for improving food quality, provided new knowledge can be transmitted (through genetic manipulation) or interpreted within the context of a working knowledge of lipid biosynthesis.

The ever increasing specialization within scientific disciplines makes it increasingly difficult to sustain an active integration between production and postproduction sciences. Disciplines focused on postproduction utilization of bioresources are typically placed in a reactionary position in dealing with issues and problems in agriculture. As a discipline, food science has historically been resigned to attempt to accommodate the compositional and associated nascent qualities of any primal food resource, with the goal of promoting, improving, or maintaining quality during any further transformation into comestibles.

The emergence of biotechnology, and specifically, genetic manipulation of organisms, provides renewed opportunities for food scientists to lend perspective toward efforts directed at genetically manipulating various traits of bioresources to
promote quality as foods or food ingredients. (With the advent of molecular biology "kits," some food scientists are positioned to bridge the production/postproduction gap as individuals.) However, with these new opportunities come responsibilities. While continuing advances in food science will expand knowledge on compositional determinants of quality of various food resources, these new developments must be interpreted in a context that will provide suitable targets for genetic manipulation. Thus, if food scientists cannot effectively position or communicate their findings in a manner conducive to exploitation by scientists in allied disciplines, the result will be manifest as lost opportunities. Specifically, opportunities exist to improve or design the nascent quality of a bioresource for the express purpose of postproduction utilization as food. This opportunity presents to us another, and perhaps the ultimate, dimension of control of food quality.

Fats and oils comprise one of the most important and abundant renewable food resources. Sources of edible fats and oils can be classified as being of plant or animal (marine and domesticated) origin. While there has been interest in the culture of microorganism for lipid production, particularly oleaginous fungi and some yeasts (1,2), one can anticipate a lengthy interim before microbial lipids contribute substantially to global supplies of edible oils. If, or when, microbial lipid production becomes commercially feasible, it would seem likely that it would gravitate to niche markets, such as the production of specialized lipids (3), including those with pharmacological properties (4). Table 1 offers a global representation of the production of edible (and selected nonedible) fats and oils (hereafter, referred to as oils or lipids) (5–7). Crude oils are refined for the purpose of purifying the triacylglycerol components for food use. These triacylglycerols serve as storage lipid (Sec. II) for the host plant tissue. Plant fruit [e.g., palm (Elaeis guineensis), coconut (Cocos nucifera),

Table 1  Global Production of Edible Oils

<table>
<thead>
<tr>
<th>Source</th>
<th>Amount (MMT)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soybean</td>
<td>18.45</td>
</tr>
<tr>
<td>Palm</td>
<td>13.96</td>
</tr>
<tr>
<td>Rapeseed</td>
<td>9.95</td>
</tr>
<tr>
<td>Sunflower</td>
<td>7.65</td>
</tr>
<tr>
<td>Tallow/grease</td>
<td>7.0</td>
</tr>
<tr>
<td>Butter</td>
<td>5.3</td>
</tr>
<tr>
<td>Groundnut</td>
<td>4.18</td>
</tr>
<tr>
<td>Cottonseed</td>
<td>3.58</td>
</tr>
<tr>
<td>Coconut</td>
<td>2.94</td>
</tr>
<tr>
<td>Palm kernel</td>
<td>1.86</td>
</tr>
<tr>
<td>Olive</td>
<td>1.85</td>
</tr>
<tr>
<td>Corn</td>
<td>1.66</td>
</tr>
<tr>
<td>Marine oils</td>
<td>1.1</td>
</tr>
<tr>
<td>Sesame</td>
<td>0.69</td>
</tr>
</tbody>
</table>

*MMT, million metric tons.
Source  Refs. 5–7.
olive (Olea europea), and avocado (Persea americana)] and seed lipids account for over 80% of world production. In the United States, plant oils account for over 94% of the annual demand of edible oils (8). Furthermore, the plant sources included in Table 1 are useful because of their suitability for organized agriculture (5) and acceptable yield of biomass (and, therefore, lipid). Less than 0.1% of about 300,000 known plant species are cultivated in organized agriculture, and fewer than 20 species account for the principal lipid products of commerce (5). However, the great diversity among the plant kingdom offers potential for exploitation in view of the current ability to genetically transfer and manipulate specific plant traits. For example, a species that produces an unusual, precious, or otherwise noteworthy fatty acid or assembled lipid but is not suitable for commercial cultivation can serve as germ plasm for incorporating these valuable traits into species that are subject to commercial culture. Thus, in view of the relative contribution to commerce and exploitable diversity of plant lipids, it seems appropriate to restrict this chapter to a discussion of storage lipid biosynthesis in plants. Reviews on fatty acid and glycerolipid biosynthesis in animal and microbial systems are available elsewhere (9–12).

II. LIPID DIVERSITY AMONG PLANTS

As diverse as lipids are in plants (and other living organisms), they can be easily classified into two categories based on their physiological role. Functional lipids are those involved in vital and metabolic processes. Typically, these are isoprenyl-derived lipids (e.g., sterols, carotenoids), and polar glycerolipids (e.g., galactosyl- and phospholipids), components that comprise, and contribute to the functioning of, (sub)cellular membranes (10,13). In view of the role of functional lipids in membranes, their fatty acid composition in plants is essentially restricted to palmitic (16:0),* palmitoleic (16:1\(_{\Delta 9}\)), trans-\(\Delta 3\)-hexadecenoic (16:1\(_{\Delta 3}\)), hexadecatrienoic (16:3\(_{\Delta 7,10,13}\)), stearic (18:0), oleic (18:1\(_{\Delta 9}\)), linoleic (18:2\(_{\Delta 9,12}\)), and linolenic (18:3\(_{\Delta 9,12,15}\)) acids (10,13,14). Functional lipids are not discussed in this chapter, except as they relate to the biosynthesis of fatty acids and the assembly of triacylglycerols (Sec. V). (Some functional lipids, such as soybean lecithin, are products of commerce, but many products of these types are simply by-products of the edible oil industry and would probably not alone justify the use of the host plant tissue in commerce.) The interested reader is encouraged to consult reviews on biosynthesis of polar lipids (15,16) and isoprenyl lipids, including sterols, carotenoids, tocopherol, and lower isoprenoids (17–19).

Storage lipids are deposited as oil bodies in fruit or seeds and are intended to serve as a carbon (energy) source for growth during germination of seeds. In fruit tissues, the raison d’être of the copious quantities of oil may be to attract predators and facilitate seed dispersal. Invariably, storage lipids predominantly take the form of triacylglycerols (20), and their fatty acyl groups are first mobilized for catabolic processes by endogenous lipases (21) and, ultimately converted into metabolic carbon (acetate) via β oxidation in peroxisomes (22). There is a much greater diversity in fatty acid composition (range of 2–24 acyl carbons) of storage triacylglycerols

*Fatty acid (or acyl group) identification is formalized throughout the chapter as X:Y, where X is the acyl carbon number and Y is the number of double bonds, all of which are cis unless otherwise noted. Subscript \(\Delta\) identifies position of the double bond, counting from the carboxy terminus.
<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>Species/source</th>
<th>Fatty acids in storage lipids (wt %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>6:0</td>
<td>Palm kernel</td>
<td>0–0.8</td>
</tr>
<tr>
<td></td>
<td>Coconut endosperm</td>
<td>0–0.6</td>
</tr>
<tr>
<td>8:0</td>
<td><em>Cuphea cyania</em> seed</td>
<td>67.8</td>
</tr>
<tr>
<td>10:0</td>
<td><em>Cuphea koehneana</em> seed</td>
<td>91.6</td>
</tr>
<tr>
<td></td>
<td><em>Cuphea lanceolata</em> seed</td>
<td>83.2</td>
</tr>
<tr>
<td></td>
<td><em>Cuphea procumbrens</em> seed</td>
<td>80.1</td>
</tr>
<tr>
<td></td>
<td>Coconut endosperm</td>
<td></td>
</tr>
<tr>
<td>12:0</td>
<td>Cinnamon (<em>Cinnamomum zeylanicum</em>) bark</td>
<td>80–90</td>
</tr>
<tr>
<td></td>
<td><em>Cuphea polymorpha</em> seed</td>
<td>80.0</td>
</tr>
<tr>
<td></td>
<td><em>Cuphea wrightii</em> seed</td>
<td>53.9</td>
</tr>
<tr>
<td></td>
<td>Coconut endosperm</td>
<td>41–56</td>
</tr>
<tr>
<td></td>
<td>Palm kernel</td>
<td>41–55</td>
</tr>
<tr>
<td></td>
<td>Babassu (<em>Orbignya martiana</em>)</td>
<td>40–55</td>
</tr>
<tr>
<td>14:0</td>
<td>Nutmeg (<em>Myristica officinalis</em>) kernel (butter)</td>
<td>60–70</td>
</tr>
<tr>
<td></td>
<td><em>Cuphea palustris</em> seed</td>
<td>63.7</td>
</tr>
<tr>
<td></td>
<td>Coconut endosperm</td>
<td>13–23</td>
</tr>
<tr>
<td></td>
<td>Palm kernel</td>
<td>14–20</td>
</tr>
<tr>
<td>16:0</td>
<td>Chinese vegetable tallow (<em>Sapium sebiferum</em>)</td>
<td>62</td>
</tr>
<tr>
<td></td>
<td>Palm fruit</td>
<td>32–59</td>
</tr>
<tr>
<td>18:0</td>
<td>Cocoa bean (butter)</td>
<td>30–36</td>
</tr>
<tr>
<td>20:0</td>
<td>Rambutan (<em>Nephelium lappaceum</em>) tallow (seed)</td>
<td>35</td>
</tr>
<tr>
<td></td>
<td>Kusum (<em>Scheichera trifuga</em>) seed</td>
<td>20–30</td>
</tr>
<tr>
<td></td>
<td><em>Sal fat</em> (<em>Shorea robusta</em>)</td>
<td>3.1–10</td>
</tr>
<tr>
<td></td>
<td>Groundnut (peanut, <em>Arachis hypogaea</em>)</td>
<td>1–3</td>
</tr>
<tr>
<td>22:0</td>
<td><em>Lophira alata</em> seed</td>
<td>15–30</td>
</tr>
<tr>
<td></td>
<td>Tamarind (<em>Tamarindus indica</em>) seed</td>
<td>12.2</td>
</tr>
<tr>
<td></td>
<td>Groundnut</td>
<td>1–5</td>
</tr>
<tr>
<td>24:0</td>
<td>Tamarind seed</td>
<td>22.3</td>
</tr>
<tr>
<td></td>
<td>Groundnut</td>
<td>0.5–3</td>
</tr>
<tr>
<td>18:1&lt;sub&gt;6&lt;/sub&gt;</td>
<td><em>Apium leptophyllum</em> seed</td>
<td>86.5</td>
</tr>
<tr>
<td></td>
<td>Coriander seed</td>
<td>75.3</td>
</tr>
<tr>
<td></td>
<td>Carrot seed</td>
<td>70–72</td>
</tr>
<tr>
<td>18:1&lt;sub&gt;12,14-OH&lt;/sub&gt;</td>
<td><em>Castor</em> bean</td>
<td>90</td>
</tr>
<tr>
<td>18:1&lt;sub&gt;12,13-epoxy&lt;/sub&gt;</td>
<td><em>Vernonia galamensis</em> seed</td>
<td>73–78</td>
</tr>
<tr>
<td></td>
<td><em>Euphorbia lagascae</em> seed</td>
<td>60</td>
</tr>
<tr>
<td>18:2&lt;sub&gt;12&lt;/sub&gt;</td>
<td><em>Safflower</em> seed</td>
<td>55–81</td>
</tr>
<tr>
<td></td>
<td><em>Sunflower</em> seed</td>
<td>20–75</td>
</tr>
<tr>
<td></td>
<td><em>Cuphea fruticosa</em> seed</td>
<td>67.2</td>
</tr>
<tr>
<td></td>
<td>Soybean</td>
<td>44–62</td>
</tr>
<tr>
<td></td>
<td>Corn kernel (germ)</td>
<td>34–62</td>
</tr>
<tr>
<td></td>
<td>Cottonseed</td>
<td>33–59</td>
</tr>
<tr>
<td>18:3&lt;sub&gt;12,15&lt;/sub&gt;</td>
<td><em>Linseed</em></td>
<td>59.8</td>
</tr>
<tr>
<td></td>
<td><em>Cuphea purpurascens</em> seed</td>
<td>30.8</td>
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<tr>
<td></td>
<td>Rapeseed (canola)</td>
<td>5–16</td>
</tr>
<tr>
<td>18:3&lt;sub&gt;6,9,12&lt;/sub&gt;</td>
<td><em>Borage</em> seed</td>
<td>19–25</td>
</tr>
<tr>
<td></td>
<td><em>Black currant</em> seed</td>
<td>15–19</td>
</tr>
<tr>
<td></td>
<td><em>Evening primrose</em> seed</td>
<td>8–14</td>
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Table 2 (Continued)

<table>
<thead>
<tr>
<th>Fatty acid&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Species/source&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Fatty acids in storage lipids (wt %)&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>20:1&lt;sub&gt;Δ5&lt;/sub&gt;</td>
<td>Meadowfoam seed</td>
<td>66.6</td>
</tr>
<tr>
<td>20:1&lt;sub&gt;Δ11&lt;/sub&gt;</td>
<td>Rapeseed (high erucic)</td>
<td>3–15</td>
</tr>
<tr>
<td>20:1&lt;sub&gt;Δ11,14-OH&lt;/sub&gt;</td>
<td><em>Lesquerella grandiflora</em> seed</td>
<td>54</td>
</tr>
<tr>
<td>22:1&lt;sub&gt;Δ13&lt;/sub&gt;</td>
<td><em>Crambe abyssinica</em> seed</td>
<td>55.7</td>
</tr>
<tr>
<td></td>
<td>Rapeseed (high erucic)</td>
<td>5–60</td>
</tr>
<tr>
<td></td>
<td><em>Mustard (Brassica carinata)</em> seed</td>
<td>22–50</td>
</tr>
<tr>
<td>22:1&lt;sub&gt;Δ5&lt;/sub&gt;</td>
<td>Meadowfoam seed</td>
<td>12</td>
</tr>
<tr>
<td>22:2&lt;sub&gt;Δ5,13&lt;/sub&gt;</td>
<td>Meadowfoam seed</td>
<td>18</td>
</tr>
<tr>
<td>24:1&lt;sub&gt;Δ5&lt;/sub&gt;</td>
<td>Honesty seed</td>
<td>24.0</td>
</tr>
</tbody>
</table>

<sup>a</sup>Fatty acid species are defined as indicated in footnote 1 in Sec. II.

<sup>b</sup>If not noted in this table, species identification appears in text upon the first mention of plant species.

<sup>c</sup>Values are either ranges or maximal values reported in the literature.


relative to functional lipids, because of the lack of any known specific structure-function role for the former in plant tissues (15,23). In addition, while functional lipids typically comprise less than 1% of weight of the plant tissue, storage triacylglycerols can reach up to 66% by weight of the oil-bearing plant tissue (20).

The great diversity in lipid composition of some oil-bearing plants is accentuated by the tendency of some species to accumulate storage triacylglycerol that is as much as 90% pure in a single fatty acid (Table 2) (24–28). Since functional and physical properties of lipids in foods are directly related to the chain length and degree of unsaturation of the constitutive fatty acids, selection of an appropriate lipid for a specific food system can be based, in part, on fatty acid composition. Thus, if the needs for a particular application of lipids in food call for properties representative of a specific fatty acid of 8–22 acyl carbons and up to three double bonds, chances are there is a plant species that deposits triacylglycerols highly enriched in that particular fatty acid. If not, or if the particular plant species does not lend itself to organized agriculture for oil production, then the trait that provides this fatty acid is subject to transfer into a species that can be commercially cultivated.

The genus *Cuphea* is of particular interest, since several species accumulate various medium chain length fatty acids in storage triacylglycerols, and these lipids are of interest from a nutritional perspective (29,30) as well as having potential to supplement or substitute for tropical oils (27). Lipids in *Cuphea* spp. range from 16% to 42% by weight of seed (27). Other plant species accumulate storage triacylglycerols enriched in fatty acids that either have unusual patterns or positions of unsaturation, or are oxygenated (20,24,26). Many of these species have potential for industrial (nonfood) uses in oleochemical industries, but others are considered to have potential applications in products that are capable of promoting health in humans (31,32), and thus, have relevance to the anticipated development of nutraceuticals. From an academic viewpoint, the genetic diversity that creates the chemotaxonomy among members of the plant kingdom, as illustrated in Table 2, provides the
tools for using genetic transformation as a means to study the process and control of lipid biosynthesis in plants. In turn, this will provide greater understanding of how to manipulate fatty acid composition in plant storage lipids for specific purposes or end uses.

It is also well known that the arrangement or distribution of fatty acyl groups along the glycerol backbone of triacylglycerols (i.e., profile of molecular species of triacylglycerols) also impacts greatly on the composite physiochemical or nutritional properties of a particular lipid (33,34). Lipids of similar fatty acid composition may have markedly different physiochemical properties because of how the fatty acids are distributed. The origins of positional distribution of fatty acids lie in triacylglycerol assembly in plants, an area in which definitive research is barely a decade old (20,35). However, a working knowledge of how triacylglycerols are specifically assembled in different species will be paramount to identifying strategies by which to modify the process in vivo and to direct triacylglycerol assembly in a manner designed to improve the functional properties of the lipid resource in foods. Section V explores this phase of lipid biosynthesis in detail.

III. FATTY ACID BIOSYNTHESIS

A. Comparative Biochemistry of Fatty Acid Synthetase Systems

1. Historical Perspective

It has been about 50 years since fatty acid synthesis was first demonstrated in cell-free extracts (36). Some of the earliest studies revealed the essential features of fatty acid biosynthesis: it required ATP, NADPH, CO$_2$ (HCO$_3^-$), CoA, Mn$^{2+}$, acetate, and source of heat-labile (FAS enzymes) and heat-stable (ultimately determined to be acyl carrier protein) proteinaceous factors isolated from the organism of interest. Slowly, each discrete, protein-associated unit activity of the type I and II fatty acid synthetase (FAS) systems was identified. Table 3 gives some of the seminal citations (37) that formalized the identification of these components and lists the unit activities (38–55). Even though virtually all the FAS enzymes, and corresponding steps in fatty acid biosynthesis, had been isolated and identified by the early 1980s, discoveries and challenges to dogmatic views regarding this intensely researched area of primary metabolism have continued. A case in point is the discovery and elucidation over the last decade of an enzyme isoform, now believed to cause the initial condensation reaction in type II FAS systems in plants and bacteria (43,44,56), and the repeated speculation that there is another condensing enzyme yet to be discovered (44,56–59).

In contrast to the wealth of history of research on fatty acid biosynthesis, definitive studies on triacylglycerol assembly and control of this process extend barely a decade into the past (15,20,35). The study of both fatty acid biosynthesis and assembly into triacylglycerols (and other glycerolipids) is essentially research in enzymology. The advent of genetic manipulation of organisms has made progress in both these areas of lipid synthesis extremely rapid since about 1990. While the theme of this chapter is founded on the characteristics of the individual enzymes involved in storage lipid biosynthesis, the superseding objective is to convey how these individual steps are coordinated as part of this overall process.
2. Basic Features

The functional units of FAS are essentially identical for all living organisms (10). These functional units will be designated by the acronyms appearing in Table 3, in boldface type, for the balance of this chapter. The distinction between FAS in various classes of organisms (plants, animals, microbes) lies in compartmentation and organizational structure of FAS, as well as in subtle differences in reaction selectivities and activities of the functional units [viz., enzymes and acyl carrier protein (ACP)] of FAS. These subtle differences account for the markedly different patterns and profiles of fatty acid biosynthesis in various classes of organisms.

Because fatty acid biosynthesis is energy intensive, it is subject to metabolic regulation and requires insulation from competing pathways or premature termination. Fatty acid biosynthesis is metabolically channeled in plants by virtue of an "ACP track" in the plastidic compartment, where the lengthening acyl chain remains covalently linked to ACP (or to the condensing enzyme) until completely synthesized. The putative product of FAS systems is palmitic acid in animals, palmitoyl-CoA in yeasts and fungi, and palmitoyl-ACP in plants and lower bacteria. Some organisms, especially oil-bearing plants, have the important capability of premature termination of fatty acids biosynthesis, giving rise to fatty acids of shorter chain lengths as the principal products (e.g., oils rich in lauric acid). The synthesized fatty acids are then exported to a "CoA track" in the endoplasmic reticulum, which serves to channel fatty acyl residues through subsequent and characteristic modification processes of oxygenation, elongation, and desaturation. Finally the fatty acids are assembled into storage and functional glycerolipids, principally in the same cellular compartment.

3. Organization and Localization

FAS systems can be classified as type I or II systems. [Historically, type III systems, or palmitate elongases, have been used to classify elongation activities yielding acyl chains longer than that of palmitate (10)]. Type I FAS systems are characterized by multifunctional protein domains, whereas type II FAS systems are characterized by dissociable proteins, each having a specific, unit activity of the FAS system.

Eukaryotes (and some higher prokaryotes, such as Mycobacterium and Corynebacterium) have type I FAS systems, as characterized from liver, adipose, and mammary and uropygial gland tissues, where the full complement of FAS activities is located on a single polypeptide domain of about 240 kDa (9). This single polypeptide has three functional subdomains, one possessing substrate entry and condensation activities of ACAT, MCAT, and KAS; the second possessing ACP and reducing activities of KHAD, EAR, and KAR; and the third, or C-terminal domain, having TE.* In situ, these FAS systems exist as homodimers where the substrate entry/condensation subdomain of one monomer channels the lengthening acyl chain to the reducing subdomain of the other monomer (i.e., functional division of dimer is not identical to the subunit division of the dimer).

Organizationally, another type I FAS system, such as in yeast, fungi and some higher bacteria, is characterized by a heteromer of two protein domains (α,β), each possessing a portion of the total FAS activities (9,60,61). The α-chain has ACP,
<table>
<thead>
<tr>
<th>Common name (other common names)</th>
<th>Abbreviation</th>
<th>EC number</th>
<th>Systematic name</th>
<th>Comment</th>
<th>Seminal citations (source organism)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyruvate dehydrogenase (pyruvate decarboxylase)</td>
<td>N/A</td>
<td>1.2.4.1</td>
<td>Pyruvate:lipoamide 2-oxidoreductase (decarboxylating and acceptor-acetylating)</td>
<td>Requires thiamine diphosphate</td>
<td>38 (animal)</td>
</tr>
<tr>
<td>Acetyl–CoA carboxylase</td>
<td>ACCase</td>
<td>6.4.1.2</td>
<td>Acetyl-CoA:carbon dioxide ligase (ADP-forming)</td>
<td>Requires biotin and ATP</td>
<td>39 (plant)</td>
</tr>
<tr>
<td>Acetyl-CoA:ACP transacylase (ACP S-acetyltransferase)</td>
<td>ACAT</td>
<td>2.3.1.38</td>
<td>Acetyl-CoA:[acyl carrier protein] S-acetyltransferase</td>
<td></td>
<td>40 (bacterial) 41 (bacterial)</td>
</tr>
<tr>
<td>Malonyl-CoA:ACP transacylase (ACP S-malonyltransferase)</td>
<td>MCAT</td>
<td>2.3.1.39</td>
<td>Malonyl-CoA:[acyl carrier protein] S-malonyltransferase</td>
<td></td>
<td>41 (bacterial) 46 (plant)</td>
</tr>
<tr>
<td>Keto-acyl-ACP synthetase (3-oxoacyl-ACP synthetase)</td>
<td>KAS (isoforms I, II, and III)</td>
<td>2.3.1.41</td>
<td>Acyl-[acyl carrier protein]:malonyl-[acyl carrier protein] C-acyltransferase (decarboxylating)</td>
<td></td>
<td>42 (bacterial) 43 (bacterial KAS III) 44 (plant KAS III)</td>
</tr>
<tr>
<td>Keto-acyl ACP reductase (3-oxoacyl-ACP reductase)</td>
<td>KAR</td>
<td>1.1.1.100</td>
<td>(3R)-3-Hydroxyacyl-[acyl carrier protein]:NADP$^+$ oxidoreductase</td>
<td>NADPH-specific</td>
<td>45 (plant)</td>
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<tr>
<td>[3-Oxoacyl-ACP synthetase (NADH)]</td>
<td>1.1.1.212</td>
<td>(3R)-3-Hydroxyacyl-[acyl carrier protein]:NAD$^+$ oxidoreductase</td>
<td>NADH-specific</td>
<td>46 (plant)</td>
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<tr>
<td>Enzyme Name</td>
<td>EC Number</td>
<td>Substrate</td>
<td>Description</td>
<td>Source</td>
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<tr>
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<td>-------------</td>
<td>--------</td>
<td></td>
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<tr>
<td>Hydroxy-Acy</td>
<td>4.2.1.58</td>
<td>(3R)-3-Hydroxybutanoyl-[acyl carrier protein] hydrolyase</td>
<td>Specific for C4:0–C8:0 derivatives</td>
<td>47 (bacterial)</td>
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<td>ACP dehydrase</td>
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<tr>
<td>(Crotonyl–ACP hydratase)</td>
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<tr>
<td>(3-Hydroxyoctanoyl–ACP dehydratase)</td>
<td>4.2.1.59</td>
<td>(3R)-3-Hydroxyoctanoyl-[acyl carrier protein] hydrolyase</td>
<td>Specific for C6:0–C12:0 derivatives</td>
<td>48 (bacterial)</td>
<td></td>
</tr>
<tr>
<td>(3-Hydroxydecanoyl–ACP dehydratase)</td>
<td>4.2.1.60</td>
<td>(3R)-3-Hydroxydecanoyl-[acyl carrier protein] hydrolyase</td>
<td>Specific for C10:0 derivatives</td>
<td>49 (yeast)</td>
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<tr>
<td>(3-Hydroxypalmitoyl–ACP dehydratase)</td>
<td>4.2.1.61</td>
<td>(3R)-3-Hydroxypalmitoyl-[acyl carrier protein] hydrolyase</td>
<td>Specific for C12:0–C16:0 (esp. C16:0) derivatives</td>
<td>48 (bacterial)</td>
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<td>Enoyl–acyl</td>
<td>1.3.1.9</td>
<td>Acyl-[acyl carrier protein]: NAD⁺ oxidoreductase</td>
<td>NADH-specific, acts on C4:0–C16:0 derivatives</td>
<td>45 (plant)</td>
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<tr>
<td>ACP reductase</td>
<td></td>
<td></td>
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<td>[Enoyl–ACP reductase (NADH)]</td>
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<tr>
<td>[Enoyl–ACP reductase (NADPH, A-specific)]</td>
<td>1.3.1.39</td>
<td>Acyl-[acyl carrier protein]: NADP⁺ oxidoreductase</td>
<td>NADPH-A-specific</td>
<td>51 (animal)</td>
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<td>Acyl–ACP hydrolase (oleoyl–ACP hydrolase; thioesterase)</td>
<td>3.1.2.14</td>
<td>Acyl-[acyl carrier protein] hydrolase</td>
<td>Selective for oleoyl-ACP</td>
<td>53, 54 (plant)</td>
<td></td>
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<tr>
<td>Acyl carrier protein</td>
<td>ACP</td>
<td>N/A</td>
<td></td>
<td>55 (plant, bacterial)</td>
<td></td>
</tr>
</tbody>
</table>

*N/A, not available or not appropriate.
*EC numbers from Ref. 37.
*Seminal citations and organism constituting source from which functional unit was definitely characterized either were taken from Ref. 37 or were among the first reports of that functional unit/isoform.

Source Ref. 37 and other works indicated.
KAS, and KAR while the β-chain has ACAT, MCAT, KHAD, EAR, and an acyl-ACP:CoA acyltransferase (instead of TE; these FAS systems produce acyl CoA instead of fatty acid as the final product of FAS).

Type II or “dissociable” FAS systems are characteristic of prokaryotes (Escherichia coli being the working example) and plants (10). The similarity of FAS in bacteria and plants is founded on the endosymbiont or “plastid–reticular” theory of evolutionary linkage between chloroplasts (and plastids in general) and the bacterial cell (62). Because of this relationship, studies on E. coli have often been viewed as a model for FAS in plants (11). Thus, some of the characteristic distinctions between E. coli and plant FAS systems are identified in this chapter because these differences help promote an understanding of the nature and control of fatty acid biosynthesis in plants.

FAS is organized in plants by virtue of being almost exclusively localized in plastidic compartments (63), although traces of FAS activity and ACP have been reported in mitochondria (64,65). Despite little evidence for the association of the unit activities of type II FAS systems as multienzyme complexes in plants in vivo (15), there are several features of control of FAS in plants (13,15,65). Unlike type I FAS systems, which exist as a single (or dual) isoform or protein domain, many of the component enzymes of type II FAS systems often exist as several isoforms, with each isoform having different properties/selectivities that affect the nature of the specific reaction they mediate. Second, many of the individual enzymes of type II FAS systems are extracted and purified as dimers and tetramers, suggesting that oligomers exist in vivo and may be more subject to control than monomeric forms. Third, the different enzymes of type II FAS systems are not present at stoichiometric levels (66,67), and individual levels and ratios of activities of the enzymes can allow for dynamic, multilocus control over fatty acid biosynthesis. The details of some of these features of control of FAS systems in plants are provided in Sec. III.C.

B. Biosynthesis of the Acyl Group and Other Plastidic Reactions

1. Basic Features

The reactions involved in synthesis of the acyl chain are shown in Figure 1. Historically, the different phases of fatty acid biosynthesis have been referred to as activation, initiation, condensation, elongation, and termination. The essential features of fatty acid synthesis are (a) initial and sustained commitment of carbon to this biosynthetic pathway, (b) utilization of acetyl and malonyl fragments as building blocks of fatty acid synthesis, and (c) lengthening of the acyl chain in two-carbon increments.

Acetyl CoA is the basic building block of fatty acid synthesis (Fig. 1). Acetate can be imported from extraplastidic sources, or alternatively, intraplastidic acetyl CoA can be formed from pyruvate by pyruvate dehydrogenase action (68). This latter pathway is likely to be of greater importance in nongreen plastids than in chloroplasts because of the more active glycolytic and amylolytic systems in the former (13,15,68–71).

2. Activation

Activation of acetate for fatty acid metabolism (either catabolic or anabolic) involves the formation of a high energy thiol ester bond, supplied by conjugation of acetate
Figure 1  Steps of fatty acid biosynthesis in plant plastids leading to genesis of the acyl chain. Individual enzymes are boxed and in **boldface**; substrates are *italicized*. Bold arrow indicates dominant pathway; dashed arrows indicate pathways of questionable importance.
with either CoA or ACP. CoA and ACP are similar in that they both possess a 4-phosphopantetheine unit that furnishes the –SH functional group. Imported acetate is first converted to acetyl-CoA by plastidic acetyl-CoA synthetase (72), the activity of which is more than sufficient to supply carbon toward fatty acid synthesis (73). Commitment of acetyl-CoA to fatty acid biosynthesis is achieved by acetyl-CoA carboxylase (ACCase), which yields malonyl CoA (Fig. 1). The only plastidic fate of malonyl-CoA is for FAS (74), although malonyl-CoA can be used for acyl chain elongation (74) and flavonoid synthesis (75,76) in extraplastidic compartments.

Plastid ACCase in plants exists in two forms: a multifunctional protein (MF-ACCase: single protein domain or “eukaryotic” form) and a multisubunit complex (MS-ACCase: “prokaryotic” form), both of which have the same three functional regions and activities relevant to fatty acid biosynthesis (13,65). These functional regions are biotin carboxyl carrier protein (BCCP), biotin carboxylase (BC), and carboxyltransferase (CT). BC requires Mg²⁺/Mn²⁺ and utilizes HCO₃⁻ and ATP to carboxylate the biotin prosthetic group of BCCP. CT then transfers the carboxyl group to acetyl-CoA to yield malonyl-CoA. The MF-ACCase also recognizes propionyl-CoA as a substrate for carboxylation (which would give rise to fatty acids of odd-numbered chain length) and has a reduced Kₘ for acetyl-CoA relative to the MS-ACCase (13). It appears that plastids of both di- and monocotyledons generally have MS-ACCase, whereas some members of the Gramineae family of plants (grasses and cereals) have MF-ACCase as the plastidic form. Although little else is known regarding the role of ACCase types in fatty acid biosynthesis, the differential organization of ACCase seems to explain the sensitivity of some Gramineae plants to grass-specific herbicides (77,78).

The next requirement for fatty acid biosynthesis is to import acetyl-CoA and malonyl-CoA groups into the FAS system via thioester exchange with ACP (Fig. 1). This exchange is achieved by the activities of acetyl–CoA:ACP transacylase (ACAT) and malonyl–CoA:ACP transacylase (MCAT), respectively (10,11,13,15). With the realization that 3-ketoacyl–ACP synthetase III (KAS III) (see Secs. III.B.3 and III.C) is primarily responsible for initiation of fatty acid biosynthesis and utilizes acetyl-CoA instead of acetyl-ACP as substrate (43,44), the exact role of ACAT has become rather nebulous. Two isoforms for both ACAT and MCAT have been reported (79,80), and for MCAT, there appears to be considerable variability of the enzyme within the plant kingdom (74). Both MCAT isoforms appear to be in leaf tissue, but only one form is dominant in seeds (80).

3. Initiation

The “initiation” step involves the conjugation of a malonyl and an acetyl group and results in the formation of the 3-ketobutanoyl–ACP (acetoacetyl-ACP) derivative with the liberation of CO₂ (Fig. 1). In both plants and E. coli, this step is now believed to be catalyzed principally by KAS III (11,43,44,81). KAS exists in two other isoforms, KAS I and KAS II, both of which are involved in chain elongation steps (Sec. III.B.5). KAS III is the most recently identified isoform, first in E. coli (43) and then in plants (44,56). KAS III is unique in that it strictly (59) utilizes acetyl-CoA (instead of acetyl-ACP) and malonyl-ACP in forming acetoacetyl-ACP. Thus, the former view (and dogma) that ACAT and acetyl-ACP levels were central to control of flux in fatty acid biosynthesis is now very doubtful (81).
The three KAS isoforms are assigned on basis of sensitivity to cerulenin and thiolactomycin (Fig. 2) (44,56,82–84), acyl-ACP chain length specificities (43, 44,66,73), and status as specific gene products of E. coli (11). KAS I is very sensitive (100% inhibition to 10 μM) to cerulenin and is the product of fabB; KAS II is moderately sensitive (50% inhibition at 50 μM) to cerulenin and is the product of fabF; KAS III is insensitive to cerulenin, but sensitive to thiolactomycin and is the product of fabH (85). Complementary DNA (cDNA) of plant KAS III is 45% identical to E. coli KAS III (FabH) and has little homology with other plant KAS isoforms (86). The three KAS isoforms have a functional cysteine at the active site, consistent with their common mechanism (85) in forming an S-acyl intermediate (and then thio exchange of the condensation product with ACP-SH to allow subsequent reactions: Sec. III.B.4) and the expected mode of action of these inhibitors (83,85,87). The differential inhibitor sensitivities have been attributed to different active site homologies and topographies, which in turn modulate the effectiveness of these −SH-directed inhibitors (87). KAS I acts on 4- to 14-carbon acyl-ACP substrates, KAS II acts only on 14- to 16-carbon acyl-ACP substrates, and both use malonyl-ACP as the condensing unit (13,36,87). KAS III principally forms acetoacetyl-ACP from malonyl-ACP and acetyl-CoA, but in isolated FAS systems where KAS I and II are both inhibited by cerulenin, acyl-ACP products of only up to four to six carbons are formed (44,87).

A KAS IV isoform in E. coli (assigned to fabJ) was inferred from genetic and cerulenin sensitivity studies (57). This isoform was also projected to be in plants to account for the observed accumulation of 8- to 10-carbon acyl-ACP in cerulenin-inhibited FAS systems. However, a subsequent report suggests that KAS IV (presumptively FabJ) was incorrectly assigned, and is the fabF gene product, or KAS II (88). More recent reports suggesting the presence of a KAS IV isoform in selected Cuphea species of plants that accumulate medium chain length fatty acids are addressed in Sec. B.5, "Elongation/Condensation."

4. Reduction Cycle

The next several reactions serve to reduce the 3-ketoacyl-ACP residue to a saturated acyl-ACP derivative and prepare the acyl chain for a repetitive cycle(s) of elongation/condensation reactions (Fig. 1). The 3-ketoacyl–ACP reductase (KAR) uses NAD(P)H to mediate the first reduction step. Two KAR isoforms exist in plants (45,46,89); the dominant form uses NADPH for reducing equivalents, whereas the other isoform uses NADH. In contrast, the FAS type II system in E. coli has only the NADPH-specific KAR (FabG) (11).

The KAR reduction step is followed by 3-hydroxyacyl–ACP dehydrase (HAD) action involving NAD(P)H to yield the 2-trans-enoyl–ACP derivative. HAD,

![cerulenin](image1) ![thiolactomycin](image2)

Figure 2  Inhibitors used to differentiate between 3-ketoacyl–ACP synthetase isoforms, I, II, and III.
isolated from spinach (Spinacia oleracea) and safflower (Carthamus tinctorius), has broad specificity toward 4- to 16-carbon acyl-ACP substrates, with maximal activity (evaluated in the reverse reaction) toward the 8-carbon acyl-ACP derivative (45,66,90). In comparison, there is some controversy regarding whether one or three isoforms of HAD (gene not yet identified) exist in the E. coli FAS system (11). In any event, HAD is not to be confused with 3-hydroxydecanoyl-ACP dehydrase (FabA) in E. coli; this latter enzyme mediates the specific step of isomerizing some of the transient 2-trans decanoyl–ACP product into the cis-3 derivative, providing for unsaturated fatty acid synthesis (vaccenic acid, 18:1\text{\textsubscript{9004}}11) in E. coli and related organisms under anaerobic conditions (11) [plant and animal systems require oxygen for fatty acid desaturation (10)].

To complete the reduction cycle, the 2-trans-enoyl–ACP reductose (EAR) uses NAD(P)H to yield the saturated acyl-ACP. Two dominant EAR isoforms exist, although four isoforms have been reported [65]. In contrast, E. coli has only one form: FabI (91). The major EAR isoform, type I, is a soluble homotetramer, appears to be present in leaf tissue and avocado fruit mesocarp, utilizes NADH, and has broad specificity toward acyl-ACP derivatives of 4 to 16 carbons, with greatest activity toward 2-hexenoyl– and 2-octenoyl–ACP derivatives (45,46,92,93). Type II EAR may be membrane-associated (65), prefers NADPH as a source of reducing power (but also uses NADPH), and exhibits a preference for enoyl-ACP of 10 or more acyl carbons (93). Both types have been isolated from castor (Ricinus communis) bean and safflower and rape (Brassica napus) seeds, hence appear to be generally present in seeds (45,66,90).

5. Elongation/Condensation

Following the reduction cycle, the saturated acyl-ACP is now ready to undergo condensation cycles initiated by KAS I and KAS II (Fig. 1). KAS I is primarily responsible for chain lengthening of 4- to 14-carbon acyl-ACP substrates, whereas KAS II is active only on 14- to 16-carbon acyl-ACP derivatives (66,73). KAS I and II isoforms have been identified in barley (Hordeum vulgare) chloroplasts, and each can exist as a homo- or heterodimer (94a). In E. coli, KAS I (FabB) is a homodimer, and KAS II (FabF) is the condensing enzyme for (un)saturated fatty acid lengthening from 16 to 18 acyl carbons (11). Each step initiated by KAS I or KAS II is followed by the reduction cycle (KAR-, HAD-, and EAR-mediated steps) to yield the corresponding saturated acyl-ACP derivative.

Recent suggestions of the presence of KAS IV isoforms in plant species that may mediate medium chain fatty acid elongation are founded on the preparation of cDNA clones from Cuphea wrightii (Kas A1), Cuphea hookeriana (Ch Kas 4), and Cuphea pulcherrima (Cpu Kas4) species (94b–d). These species also contain medium chain TE and accumulate medium chain fatty acids in seed oils. The deduced sequences of these cDNA clones are 98% identical to each other, about 60% identical to KAS I, and 75–85% identical to KAS II. Expression of these KAS isoforms in Brassica species leads to more than fourfold enhanced rates of extension of 6:0- and 8:0-ACP over nontransformed plants in cerulenin-inhibited seed extracts. Furthermore, coexpression of Kas4 and the gene encoding for the medium chain TE in Brassica plants gives rise to medium fatty acids in seed oils whereas none were produced from wild-type plants. Although the KAS IV has not yet been isolated, current thinking is that it is a condensing/elongation enzyme specific for medium
chain length acyl-ACP residues, and its presence is important (if not essential) to seed oils that accumulate medium chain fatty acids. See also Sec. C.2. "Flux and Feedback Control."

The putative product of plant FAS is palmitoyl-ACP (16:0-ACP). Thus, the action of KAS II represents the first elongation step [historically referred to as type III FAS (10)] that can occur, forming stearoyl-ACP (18:0-ACP) (Fig. 3). Since there is usually very little (<5%) 18:0 in plant oils (20,28), further modification steps, desaturation and/or further elongation, account for the fate and rather transient nature of 18:0.

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**Figure 3** Steps of fatty acid biosynthesis in plant plastids involved in desaturation, elongation, termination, or acyl transfer reactions. Fatty acyl groups are designated as in footnote 1. Other details are the same as in Figure 1.
6. Desaturation

The dominant fatty acid to be exported from the plastid in plants is oleic acid (18:1\(^{9}\)). It is formed by the action of \textit{Δ9}-desaturase (\textit{Δ9DES}) on 18:0-\textit{ACP} (Table 3 and Fig. 3). The \textit{Δ9DES} is a soluble, homodimeric diiron-oxo enzyme localized in the stroma of the plastid (52,95,96). The soluble nature of \textit{Δ9DES} is a unique feature of this plant enzyme relative to desaturases in general in higher organisms (10). Desaturation is mediated through electron transfer from NADPH (or reducing equivalents from PS I/II in chloroplasts) \(→\) ferredoxin-NADPH reductase \(→\) ferredoxin \(→\) desaturase + O\(_2\) (10,93). Although the \textit{Δ9DES} is 100 times more active toward 18:0-\textit{ACP} than 16:0-\textit{ACP}, it will also act on 14:0- and 16:0-\textit{ACP}, to yield myristoleic (14:1\(^{9}\))- and palmitoleic (16:1\(^{9}\))-\textit{ACP} derivatives.

Two isoforms have been identified in sesame (\textit{Sesamum indicum}), with one form restricted to seeds (97). Seeds of some species of Araliaceae, Garryaceae, and Umbelliferae, such as coriander (\textit{Coriandrum sativum}), dill (\textit{Anethum graveolens}), parsley (\textit{Phytophthora megasperma}), and carrot (\textit{Daucus carota}), are known to yield oils unusually rich in petroselenic acid (18:1\(^{6}\)) (Table 2 and Refs. 24 and 25). This can be explained by the presence of \textit{Δ4 acyl-ACP desaturase} (\textit{Δ4DES}) (98,99). The \textit{Δ4DES} is believed to be a stromal enzyme (based on homology to the \textit{Δ9DES}) that acts principally on 16:0-\textit{ACP}. Subsequent elongation by \textit{KAS II} (or possibly, an unidentified \textit{KAS} isoform) gives rise to 18:1\(^{6}\). Conversion of 18:1\(^{6}\) by a cleavage process would provide an industrial source of lauric and adipic acids.

7. Termination

Termination reactions are mediated by thioesterase (oleoyl-\textit{ACP} hydrolase or \textit{TE}) or \textit{acyl-ACP acyltransferase} activities (Fig. 3).

\textbf{a. Acyl-ACP Hydrolysis.} Thioesterases (\textit{TEs}) yield the corresponding fatty acid and \textit{ACP}, and the dominant \textit{TE} isoform in plants exhibits a 5- to 10-fold preference for 18:1\(^{9}\)-\textit{ACP} over 16:0- and 18:0-\textit{ACP} (53,54,95). This \textit{TE} isoform is required of all plants to provide a “housekeeping” function (100) and to channel 18:1\(^{9}\) toward further desaturation processes for incorporation into polar glycerolipids to serve as functional units of cellular membranes (13,15,100). This specificity accounts for the preponderance of 18:1\(^{9}\), and combined with the \textit{Δ9DES}, a lack of 18:0 in plant lipids. However, this \textit{TE} does not account for the presence of shorter chain length fatty acids that exist in oils of tissues of coconut, palm kernel, and \textit{Cuphea} spp.

The existence in plants of medium chain \textit{TE} isoforms has long been speculated (74), partly on the basis of an extrapolation of a medium chain \textit{TE} (and transcylase) activity in lactating mammary gland tissues (101). However, the existence of medium chain \textit{TE} isoforms in plants has been confirmed only in the last 5 years (102–104), and the currently recognized diversity of \textit{TE} isoforms in plant FAS systems may be one of the most exploitable traits. Some \textit{TE} isoforms, such as that from California bay [\textit{Umbellularia californica} (102,103), \textit{Cuphea wrightii} (58), camphor (\textit{Cinnamomum camphora}), and elm (\textit{Ulmus americana}) (104), are selective for hydrolyzing 10:0- and 12:0-\textit{ACP}, resulting in accumulation of oils rich in lauric acid. The \textit{TE} isoform(s) in coconut is more broadly selective for 8- to 12-carbon acyl-\textit{ACP} substrates (104), and two \textit{TE} isoforms from \textit{Cuphea palustris} have selectivities toward...
b. Acyl Transfer to Plastidic Glycerol-3-phosphate. The other possible fate of neosynthesized acyl groups by plastidic FAS is the incorporation into functional glycerolipids in the plastid. **Acyl–ACP acyltransferases** exist in plastids and include acyl-ACP: *sn*-glycerol-3-phosphate acyltransferase (GPAT) and acyl-ACP: 1-acyl-*sn*-glycerol-3-phosphate acyltransferase (or lysophosphatidic acid acyltransferase, LPAAT) (Fig. 3 and Refs. 15 and 35). These two enzymes serve to channel newly synthesized acyl groups toward polar glycerolipid synthesis within the plastid (principally for phosphatidylglycerol, galactoglycerolipid, and sulfoglycerolipid assembly) (15,16). These plastidic acyltransferases are in competition with **TE**, which mediates the release of fatty acids for export to the endoplasmic reticulum for further processing. The relative activities and selectivities of plastidic acyl–ACP acyltransferases, TE, Δ9DES, and KAS II provide the basis for partitioning of fatty acids among competing processes or fates (Fig. 3).

The pattern of fatty acid flux that occurs provides the basis for a chemotaxonomic classification of plants as “prokaryotic” or “16:3” plants, or alternatively, “eukaryotic” or “18:3” plants, based on the enrichment of 16-carbon acyl residues at the *sn*-2 position for the former class (106,107). Plants having physiologically relevant plastidic acyl–ACP acyltransferase activities and a specific ACP isofrom (see Sec. III.C.4) yield the “prokaryotic” configuration among some of the plastidic polar glycerolipids and are characterized as “16:3” species. Since plastidic acyltransferases have little to do with triacylglycerol synthesis and oil deposition, they are not considered in much detail here and the reader is referred to reviews covering this topic (15,106,108). However, plastidic acyltransferases are identical (same EC numbers: see Ref. 37) to those involved in triacylglycerol assembly (see below, Sec. V.B, Table 5), except that the plastidic forms recognize acyl-ACP, but not acyl-CoA substrates (35).

The relevance of plastidic acyltransferase to FAS and triacylglycerol assembly lies in their impact on the profile of fatty acids that are exported from the plastid for triacylglycerol assembly in the endoplasmic reticulum, and in one case, they have been subject to genetic transformation in a manner that has the potential to improve resistance of chilling-sensitive plants (Sec. VI). In addition, the apparent strict specificity of the plastidic GPAT to recognize the Δ9 double bond (105) helps ensure that unusual fatty acids, like petroselenic acid (18:1Δ9), are channeled toward triacylglycerol assembly in the endoplasmic reticulum, and not toward plastidic polar glycerolipid assembly. This feature maintains the profile of unsaturated 16- to 18-carbon fatty acids in membrane lipids to allow for normal physiological functioning (13,15,23).

8. Fate of Plastidic 16:0- (18:0-) and 18:1Δ9-ACP

Although 16- to 18-acyl carbon fatty acids are the principal products of plant FAS, the diversity and abundance of fatty acid chain lengths of 8–24 among storage lipids indicates that some plant species are distinguished by premature termination of fatty acid synthesis or postsynthetic modification.
As a result of termination reactions by TE or acyl–ACP acyltransferase action, the acyl group exits the “ACP track” and ACP remains inside the plastid to be recycled for further fatty acid biosynthesis. If not mobilized for plastidic polar glycerolipid assembly, newly synthesized fatty acid is exported to the cytosol and converted to CoA derivatives during passage through the outer envelope of the plastid. Trafficking of fatty acyl-CoAs to other cellular compartments determines their fate. Import to peroxisomes, especially glyoxysomes, makes fatty acyl-CoAs subject to β oxidation (22), which would represent a futile cycle if the fatty acid was just synthesized. In the context of glycerolipid biosynthesis, the destination of the fatty acyl-CoAs would be the endoplasmic reticulum, where acyl chain modification and elongation reactions (Sec. IV) takes place, as well as triacylglycerol assembly (Sec. V).

C. Control Points of Fatty Acid Biosynthesis in Plants

Physiological points of control of metabolic processes can be generally delineated as coarse (changes in enzyme levels) or fine (changes in metabolite or effector levels). For plant FAS systems, current views of regulation are perhaps best subordinated in terms of rate-limiting step(s), flux and feedback control, developmental control, and isoform influence. There has also been much speculation that the steady-state levels of acetyl-CoA, malonyl-CoA, ACP, ATP, and nucleotides may have influence on the control of fatty acid biosynthesis in plants (74). However, aside from measurement of ACP and acyl-ACP levels in tissues during light–dark cycles to investigate regulatory control of ACCase (see next paragraph), little definitive work has been done on evaluating the role of metabolite levels. Furthermore, the presence of multiple pathways of acetyl-CoA metabolism would make it a formidable challenge to identify the features that may regulate fatty acid biosynthesis (74).

1. Rate-Limiting Step(s)

ACCase is considered to be the rate-limiting and carbon-committing step for fatty acid synthesis (11,15,109) and is subject to regulatory control. A central and unresolved issue regarding coarse control of ACCase is the uncertainty of the physiological roles of the MF-ACCase and MS-ACCase in the plant cell. Historically, it is interesting that over the past decades, the dogmatic view of the nature of ACCase has shifted. First it was thought to be strictly a dissociable MS-ACCase form (cf. E. coli ACCase), then strictly an MF-ACCase [based on the conclusion that proteolysis during isolation was responsible for dissociation into subunits (15,74)]. The current view is that both forms exist (11,110). The MF-ACCase appears to account for only about 20% of the total ACCase activity in plants and is enriched in epidermal tissues (77). Thus, the central role of MF-ACCase may be in fatty acid elongation reactions that are extraplastidic (11,110,111).

Different features exits for genomic control of these ACCase isoforms. The MF-ACCase is a single-gene product, whereas the MS-ACCase is a multiple-gene product with BCCP being nuclear-encoded, and one of the CT subunits being plastidial-encoded (11). Diurnal (light–dark) control of ACCase activity in chloroplasts has been known for some time, inasmuch as activity (of this enzyme) is suppressed in the dark (112). Subsequent studies (113,114) of acyl-ACP pools, indicating an increase in acetyl-ACP during the dark cycle, lend further support to ACCase as the rate-limiting step of fatty acid biosynthesis.
In developing castor seed, BC and ACCase expression are well coordinated, and BC increases to 0.8% of total seed protein, compared to less than 0.05% in young leaf or root tissue (110,115). This developmental signal to induce rapid deposition of storage triacylglycerols in maturing seed suggests that the rate of lipid synthesis is controlled by ACCase. In this case, the MS-ACCase is the key isoform, and BC is found throughout embryo development, whereas the MF-ACCase was detected only in early stages of embryo development.

Even though MS-ACCase is dissociable, there is evidence that complexes exist in situ. In preparing pea (Pisum sativum) chloroplasts, BC copurifies with ACCase (110), and antibodies to one of the CT subunits inhibit ACCase and interact with BCCP (116). In pea leaves, two isoforms of CT have been identified and the α-CT isoform is suggested to be membrane-associated (117). Similarly, in E. coli, homodimeric BC copurifies with homodimeric BCCP, whereas CT exists as a heterotetramer of isoforms (αβ2) (11). In terms of possible fine control mechanisms, castor and pea ACCases are not inhibited by long chain acyl-ACP, fatty acids, or ACP (110).

Greater understanding of in vivo regulation of ACCase will be possible when studies are done with transformants that have antisense constructs to ACCase or overexpress ACCase activity (118). Already there are indications of complex genomic control of ACCase. In E. coli transformants expressing a medium chain length–specific TE, increased levels of BCCP occurred, and this finding was related to enhanced ACCase activity and carbon flux through FAS (119). Despite the preponderance of evidence implicating ACCase as a key control point in FAS, it is suggested that ACCase be considered as just one of multiple control points in fatty acid biosynthesis in plants (120). For example, the examination of whether ACP levels influence the rate of fatty acid biosynthesis and storage lipid deposition has been suggested (16).

As a side note, the down-regulation of the plastidic (MS-)ACCase form is viewed as a potential means to enhance production of the biodegradable polymer polyhydroxybutyrate (121), by diverting carbon and energy resources away from common storage lipid accumulation (111). Such polymers may have impact on food packaging in the future.

ACAT was once thought to be rate-limiting to fatty acid synthesis, on the basis of a comparison of in vitro activities of isolated components of type II FAS from plant tissues (66,67). Compared to acetyl-CoA, 4:0-ACP, and 6:0-ACP, however, acetyl-ACP is the least effective primer for fatty acid synthesis (122) and is now judged to be a minor participant in FAS in plants (11,13,65); ACAT may simply provide for a backup mechanism for KAS III in FAS (65). Although ACAT activity can be resolved from KAS III activity (118), it has been suggested that ACAT activity may reside with KAS III (11), or that ACAT and KAS III are structurally related, including the possibility that ACAT is a proteolytic degradation product of KAS III (118). Finally, it has been suggested that ACAT may be an artifact derived from acyl–ACP synthetase activity that is important to phosphatidylethanolamine metabolism, extrapolating from E. coli (118). Again, the importance of ACAT likely awaits resolution from studies evaluating genetic constructs with overexpressed and suppressed levels of ACAT. One interesting observation made with ACAT is that when added to a crude FAS system, the proportion of 10:0 to 12:0 fatty acids increases, suggesting that ACAT activity can modulate the product profile of FAS in
plants (79). This observation is similar to those that form the basis of speculation that a KAS IV isoform exists in plants.

2. Flux and Feedback Control

Since KAS III causes the initial condensation step in fatty acid biosynthesis, some features of the control of this enzyme have been evaluated. ACP and acyl-ACP levels have impact on KAS III activity in vitro. Feedback inhibition of acyl-ACP, especially 10:0-ACP, of KAS III occurs in Cuphea lanceolata, a species that accumulates decanoylglycerides (59); KAS III from rape and spinach behaves similarly. Feedback inhibition of KAS III also occurs in E. coli, with increased inhibition with increasing acyl chain length, leading to an accumulation of malonyl-ACP (123); however, free ACP and fatty acids are not inhibitory.

Low levels of ACP inhibit KAS III in C. lanceolata (124), indicating a need for FAS to "budget" available ACP between initiation reactions and ongoing chain lengthening. Relatedly, overexpression of E. coli KAS III in E. coli and rape results in an increase in 14:0 and corresponding decrease in 18:1ω9 (120). This may result from a shift toward overcommitment of ACP to initiating chain synthesis and greater opportunity for premature chain termination by TE action. In accord with this interpretation, overexpressed TE leads to uncontrolled fatty acid biosynthesis in E. coli (125), where elevated steady-state ACP levels would be expected. Overexpression of plant medium chain TE in E. coli accelerates fatty acid biosynthesis with attendant increases in steady-state malonyl-ACP and fatty acid levels, mediated by rapid removal of long chain acyl-ACP (119). A cDNA clone encoding for KAS III from Cuphea wrightii should allow for further evaluation of FAS control in rapidly developing embryos of this organism (126).

Despite a recent assertion (57) and subsequent rebuttal (88) regarding the presence of a KAS IV in E. coli, several studies have led to the speculation that a KAS IV does exist in plants (and thus, likely in E. coli) (56, 59, 124). This postulation is based on the formation of acyl-ACP products having 6–10 acyl carbons in cerulenin-inhibited preparations (where only KAS III, and not KAS I/II, can act). Since KAS III specificity prohibits formation of acyl chain lengths beyond 4:0, another KAS isoform is suggested to account for the observed condensation steps beyond butyrate in cerulenin-inhibited FAS systems. In fact, recent reports (discussed in Sec. B.5) have suggested the presence of KAS IV in Cuphea spp. (94b–d). KAS IV appears to have a regulatory role in fatty acid biosynthesis in that transformed plants expressing KAS IV and medium chain TE accumulate medium chain length fatty acids in oils to a greater extent than seed oil plants expressing this TE alone. This provides evidence for a "flux-forward" regulatory aspect of KAS IV in that it could enhance steady-state levels of medium chain ACP species and provide greater opportunity for medium chain TE action to terminate elongation and allow accumulation of medium chain fatty acids in the oils.

Although ACCase in plants is not subject to the same mechanisms of control as in animal FAS systems, it may be subject to feedback inhibition (13).

3. Developmental Control on FAS

Developmental control of fatty acid biosynthesis is responsible for redirecting plant metabolism toward a rapid deposition of storage lipid in maturing seed and fruit tissues. Central to developmental control is the genomic expression of organ-specific
(viz., seed, fruit) genes, and this is a pivotal issue regarding the commercial exploitation of specific traits by genetic manipulation (Sec. VI and Chapter 28). This is because the objective of genetic transformation is to manipulate the nature of lipid accumulation in specific (storage) organs without compromising the vitality of the whole plant.

Many enzymes of FAS are known to be subject to developmental control. KAS I, II, and III are induced, in a coordination fashion, prior to lipid accumulation in developing rape seeds (42,127,128). Similarly, both Δ9DES isoforms accumulate in developing sesame seeds (97). The TE isoforms selective for medium chain acyl-ACP s in Cuphea palustris seeds are seed-specific and are developmentally regulated to coincide with seed maturation (129). BC and ACCase expression are well coordinated in developing castor seed, and it is the MS-ACCase (and not MF-ACCase) that is expressed in a manner paralleling seed development and lipid deposition (110,115). EAR is developmentally regulated and coordinated with seed embryo development and oil deposition such that activity in developing oilseeds is 15–30 times that in leaf tissues (130–132). Both EAR isoforms are highly conserved and are present in seed and leaf tissue; there is no specific seed gene for EAR (133,134).

The importance of organ-specific gene expression is exemplified by experiments with rape transformed with a medium chain–specific TE from California bay (135). Chloroplasts from both leaf and seed tissue of transformants produced 12:0 as an end product of FAS, indicating that the TE was expressed in both organs. However, in intact leaf tissue no 12:0 accumulated, indicating that newly synthesized laurate was immediately exported to the peroxisomes for β oxidation to prevent incorporation of this fatty acid in functional glycerolipids. Although this illustrates an important survival mechanism for the plant, such a futile cycle represents a waste of cellular energy.

4. Presence of Isoforms of Units of FAS in Plants

As pointed out in the preceding sections (II.A.3 and III.B), many of the component enzymes/proteins of FAS systems in plants have multiple isoforms. Substrate selectivity features are known for some of these isoforms; however, generally little is known about their influence on regulatory control of fatty acid biosynthesis.

Enzymes involved in the reduction cycle have not been considered much in this context. However, now that the gene for the minor KAR isoform has been isolated from Cuphea lanceolata (136), an understanding of a possible role for this isoform in control of fatty acid biosynthesis may develop in the next several years.

MCAT exists as at least two isoforms, and it exhibits considerable variability among plants in molecular size and kinetics (74,80), indicating some degree of regulatory control or species-specific adaptation. The presence of two isoforms each of Δ9DES (97) and a medium chain length-specific TE (129) leads us to infer some regulatory function of these specific enzymes.

Perhaps the one clear role of isoforms of FAS units resides with the function of ACP. ACP was once characterized as the heat-stable protein factor required by FAS (36,65). It exists in as many as three isoforms (137), although two isoforms appear to be dominant in plant tissues (74,138,139). Surprisingly (65), only one isoform appears to exist in seed tissue (where a greater demand for functional diversity might be expected), whereas both major isoforms are found in leaf tissue (140). ACP levels in rape and soybean (Glycine max) seed increase just prior to
deposition of storage lipids, indicating developmental control (138,139), similar to control mechanisms for other units of FAS discussed in Sec. III.C.3. Another regulatory feature of ACP action in *E. coli* is the presence of membrane-binding sites (141), although this issue has not received much attention in plant FAS systems.

The significance of ACP isoforms was perhaps best illustrated (139) by their influence on trafficking of 18:1-ACP residues toward the competing pathways of plastidic glycerolipid synthesis and export to the endoplasmic reticulum (cf. Fig. 3). In spinach leaf, acyl-ACP isoform I is a better substrate than acyl-ACP isoform II for TE, and the reverse is true for plastidic GPAT activities. Thus, ACP isoform I tends to favor channeling newly synthesized fatty acids to the endoplasmic reticulum (for further elongation, desaturation, hydroxylation, or glycerolipid assembly in the “eukaryotic” configuration), whereas ACP isoform II favors intraplastidic utilization of newly synthesized fatty acids for glycerolipid synthesis, giving rise to the “prokaryotic” configuration of plastid-derived polar glycerolipids. Despite these differences, the two ACP isoforms are equally supportive of plant fatty acid biosynthesis in general and yield similar fatty acid profiles form plant FAS systems.

### IV. EXTRAPLASTIDIC MODIFICATION REACTIONS

With reference to triacylglycerol-accumulating plant tissues, 16:0 and 18:1,9 are principal products of plant FAS systems. To account for the diversity among the 16–24 acyl carbon derivatives that can be found in seed/fruit oils, modification reactions must take place, and these occur in the endoplasmic reticulum. (Fatty acids of fewer than 18 acyl carbons exported from the plastid generally become incorporated into triacylglycerols without further modification in the endoplasmic reticulum.) The basic modification processes are oxygenation (hydroxylation, epoxidation), elongation, and desaturation, and these processes give rise to many unusual fatty acids (some of which appear in Table 2). The unusual fatty acids (>200) that exist in plants are generally confined to storage lipids (142), in keeping with the strict requirements that functional lipids have for constitutive fatty acids (13,15,23). Most of these unusual fatty acids are found at trace levels and are potential targets for use in the oleochemical industries. Later in the chapter we consider a few such modification processes (only those that yield fatty acid derivatives relevant to use in, or as, foods). Figure 4 provides an overview of common fatty acid modification reactions taking place in the endoplasmic reticulum, and the enzymes associated with these reactions are listed in Table 4.

#### A. Oxygenation

Hydroxylation of fatty acids yields derivatives that can have fragrance/flavoring potential under conditions favoring the induction of lactonization. In addition, many hydroxylated fatty acid derivatives are components of natural waxes, which are effective moisture barriers when used in foods. The most common hydroxylated fatty acid is ricinoleic acid (18:1,9,12-OH), which is abundant in the developing endosperm of castor bean. The oleate hydroxylase (12-MO) is localized in the endoplasmic reticulum; it utilizes 18:1,9-phosphatidylcholine, oxygen, and NAD(P)H channeled through cytochrome *b*₅, and it is specific for both chain length and double-bond position (143,144). Efforts to purify the 12-MO have not been successful because
of its relative lability (perhaps due to instability when removed from the membrane), but the 12-MO is believed to function in a manner similar to membrane desaturases (145,146). Indeed, the electron transfer components of the hydroxylase and membrane desaturases are identical (cf. Sec. IV.C). The reaction utilizes reducing equivalents from NADH, and hydroxylation of 18:1_9004_9 occurs only upon esterification at the sn-2 position of phosphatidylcholine (PC) (147). In spite of the lability of the

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Table 4  Enzyme Activities Involved in Fatty Acid Modification in the Endoplasmic Reticulum

<table>
<thead>
<tr>
<th>Common name (Other common names)</th>
<th>Abbreviation</th>
<th>EC number&lt;sup&gt;a,b&lt;/sup&gt;</th>
<th>Systematic name</th>
<th>Relevant citations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acyl–CoA elongase (Icosanoyl–CoA synthetase)</td>
<td>ACE</td>
<td>2.3.1.119</td>
<td>Stearoyl-CoA malonyl-CoA C-acyltransferase (decarboxylating, oxoacyl- and enoyl-reducing)</td>
<td>150 151</td>
</tr>
<tr>
<td>Oleate hydroxylase (Phosphatidylcholine 12-monoxygenase; ricinoleic acid synthase; oleate Δ&lt;sup&gt;12&lt;/sup&gt;-hydroxylase)</td>
<td>12-MO</td>
<td>1.14.13.26</td>
<td>1-Acyl-2-oleoyl-sn-glycerol-3-phosphocholine:NADH:oxygen oxidoreductase (12-hydroxylating)</td>
<td>143 144 147</td>
</tr>
<tr>
<td>Oleate epoxidase</td>
<td>12-EPOX</td>
<td>N/A</td>
<td>N/A</td>
<td>149</td>
</tr>
<tr>
<td>Oleoyl–phosphatidylcholine desaturase (Phosphatidylcholine desaturase; oleate desaturase; linoleate synthase)</td>
<td>Δ12DES</td>
<td>1.3.1.35</td>
<td>1-Acyl-2-oleoyl-sn-glycero-3-phosphocholine: NAD&lt;sup&gt;+&lt;/sup&gt;-Δ&lt;sup&gt;12&lt;/sup&gt;-oxidoreductase</td>
<td>160 161</td>
</tr>
<tr>
<td>Linoleoyl–phosphatidylcholine desaturase (α-linolenate-forming)</td>
<td>Δ15DES</td>
<td>N/A</td>
<td>N/A</td>
<td>166</td>
</tr>
<tr>
<td>Linoleoyl–phosphatidylcholine desaturase (γ-linolenate-forming)</td>
<td>Δ6DES</td>
<td>N/A</td>
<td>N/A</td>
<td>167 168</td>
</tr>
</tbody>
</table>

<sup>a</sup>EC numbers from Ref. 37.

<sup>b</sup>N/A, not available or not appropriate.

Source  Ref. 37 and other works indicated.
**12-MO**, the cDNA for the enzyme has been isolated (109) and rapeseed transformants have been prepared for field evaluation (148).

Fatty acid oxygenating and desaturating enzymes in plants appear to share similar mechanisms. The oleate epoxidase (12-EPOX) (in *Euphorbia lagascae* and, likely, *Vernonia* spp.) causes epoxidation of 18:1\(_{9}\) to yield vernolic acid (18:1\(_{9,12/13-epoxy}\)) and requires NADPH and PC-linked 18:1\(_{9}\) (149). Furthermore, antibodies to cytochrome b\(_5\) inhibited both epoxidation and desaturation (see Sec. IV.C) reactions with PC-linked 18:1\(_{9}\).

### B. Elongation

Chain elongation steps are responsible for the production of the series of gondoic (20:1\(_{11}\)), erucic (22:1\(_{13}\)), and nervonic (24:1\(_{15}\)) acids in selected *Brassica*, *Crambe*, and related species, in some of the same fatty acids plus 20:1\(_{5}\) and 22:2\(_{5,13}\) in meadowfoam (*Limnanthes alba*), in the series of arachidic (20:0), behenic (22:0), and lignoceric (24:0) acids in groundnut (*Arachis hypogaea*) oils, and in other saturated and monounsaturated long chain fatty acids that become incorporated as surface waxes (28). Very long chain fatty acid (VLCFA) synthesis has been primarily studied in seed of rape, meadowfoam, and jojoba (*Simmondsia chinesis*), and in epidermal tissue of leek (*Allium porrum*) (150). Generally, only saturated and monounsaturated acyl-CoA derivatives are substrates for elongation in seeds. The basic process of elongation (of 18:0 or 18:1\(_{9}\)) involves successive two carbon additions supplied by malonyl-CoA to the acyl-CoA chain in the presence of NAD(P)H and membraneous acyl-CoA elongase (ACE) enzymes (Table 4 and Fig. 4) (151–154).

The jojoba elongation system is distinct in that it recognizes both ACP and CoA thioesters of the lengthening acyl chain as substrate (150). For surface wax formation, the subsequent actions of a reductase and a ligating enzyme are required, as is a means of transporting the acyl chains to the cellular surface for wax deposition (146,150).

It is not certain whether a single or multiple ACE(s) is/are involved in lengthening 18:X to 24:X, but the various elongation steps appear to share common functional units (152). Studies with *Arabidopsis* mutants revealed that the product of the FAE1 (fatty acid elongase) gene was a synthetase/condensing enzyme, based on its homology with KAS III and *E. coli* (155). Furthermore, this ACE was concluded to be singularly capable of both condensation steps from 18 to 22 acyl carbons. Similar to FAS systems, the complete reduction cycle and intermediate steps are believed to occur after each condensation step (150). Indeed, a partially purified ACE system possessed the full complement of expected condensing and reducing activities (156) (as shown in Fig. 1 for FAS). The emerging organizational structure of ACE systems is based on multienzyme membrane complexes held by strong interactions with the acyl-CoA substrates partitioning at the interface to gain access to the elongase (150). The ACE in *Arabidopsis*, which is expressed in developing seeds and not leaves (155), is likely modulated by the surrounding membrane environment. At least two contentious issues remain regarding the nature of ACE action. It has been questioned whether acyl-CoA are the putative substrates, and instead, lipid-linked acyl groups or free acids have been suggested as possible substrates for elongation in oil bodies of rape (157). A second and, perhaps, related issue is the ambiguity surrounding whether a covalent acyl-enzyme intermediate is involved in the reaction...
mechanism. The absence of such an intermediate has been suggested (150), but there is also evidence of involvement of a thioester intermediate (156); the latter possibility is favored by the homology of the ACE to KAS III (155).

C. Desaturation

The initial desaturation step to form 18:1,9 is mediated by the soluble (stromal) Δ9DES of the plastid, while the acyl chain is on the “ACP track” (Fig. 3 and Sec. III.B.6). Both the plastid and the endoplasmic reticulum possess membrane desaturase systems that yield the di- and triunsaturated fatty acids characteristic of plants. Both membrane systems use glycerolipid-linked fatty acyl groups as substrates [an exception is found in meadowfoam seeds, which also have an acyl-CoA desaturase (15)]. The plastidic system acts on acyl groups linked to galactoglycerolipids, sulfoglycerolipids, and phosphatidylglycerol as substrates and gives rise to membrane lipids assembled in a “prokaryotic” configuration (see Sec. III.B.7). Since these glycerolipids have little impact on triacylglycerol composition, the plastid desaturase systems are not considered further.

The endoplasmic reticulum system almost exclusively uses phosphatidylcholine (PC) as the carrier for 18:1,9 desaturation to form 18:2,9,12 and then 18:3,9,12,15 (Fig. 4). In oil-bearing seeds and fruit, a major fraction of these polyunsaturated fatty acids becomes incorporated into storage triacylglycerols. Accordingly, the balance of this section is devoted to the endoplasmic reticular desaturases. Much of the understanding of the molecular biology of membrane desaturases has been derived from work on mutants of the small mustard plant Arabidopsis thaliana (15). This organism is suitable for genetic studies of plants because it has a relatively small genome and mutants are easy to prepare; moreover, it is phylogenetically related to Brassica species, such as rape (148).

The two membrane desaturases of the endoplasmic reticulum are defined as products of the fad2 and fad3 genes (15). Mobilization for desaturation of 18:1,9-CoA, imported from the plastid, requires the sequential action of GPAT and lyso-phosphatidic acid acyltransferase (LPAAT) to form the 18:1,9-glycerol esters, and then phosphatidic acid phosphatase (PTP) and CDP-choline:diacylglycerol cholinephosphotransferase (CPT) to yield PC-linked 18:1,9 (20) (Fig. 4; these enzymes are described in more detail in Sec. V, Table 5 and Fig. 5). The natures of GPAT and LPAAT give rise to the “eukaryotic” configuration of glycerolipids in that saturated units of 16 or 18 acyl carbons are virtually excluded from the sn-2 position in favor of 18:1,9, which is positioned at both sn-1,2 sites (158).

Since these membrane desaturases have been difficult to isolate (13,14,159), much of the understanding of their properties has been based on the behavior of microsomal preparations (14,160–162). Evidence strongly weighs in favor of multiple desaturases, each specific for a particular site (Δ6, Δ12, Δ15), rather than one desaturase capable of successive desaturation steps (14,146) (Fig. 4). The desaturases are iron-containing enzymes that require O2 as cosubstrate (terminal electron acceptor) and reducing equivalents transferred from NAD(P)H, likely through cytochromes (b5 in particular) and cytochrome flavoprotein reductases (14,161,163). Sequence analyses have revealed histidine-rich motifs, which could serve as the iron-coordinating region of the protein (164).

For the 18:1-PC Δ12 desaturase, (Δ12DES) both sn-1,2-18:1,9 residues are reactive, with the sn-2 residue more favorable for reaction (160,161). This pattern
**Table 5**  Enzyme Activities Involved in Triacylglycerol Assembly in the Endoplasmic Reticulum

<table>
<thead>
<tr>
<th>Common name (Other common names)</th>
<th>Abbreviation</th>
<th>EC number$^a$</th>
<th>Systematic name</th>
<th>Relevant citations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycerol-3-phosphate acyltransferase (Glycerol-3-phosphate O-acyltransferase)</td>
<td>GPAT</td>
<td>2.3.1.15</td>
<td>Acyl-CoA:sn-glycerol-3-phosphate 1-O-acyltransferase</td>
<td>170</td>
</tr>
<tr>
<td>Lyso phosphatidic acid acyltransferase (1-Acylglycerol-3-phosphate O-acyltransferase)</td>
<td>LPAAT</td>
<td>2.3.1.51</td>
<td>Acyl-CoA:1-acyl-sn-glycerol-3-phosphate 2-O-acyltransferase</td>
<td>171</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>172</td>
</tr>
<tr>
<td>Phosphatidic acid phosphatase (Phosphatidate phosphatase)</td>
<td>PTAP</td>
<td>3.1.3.4</td>
<td>3-sn-Phosphatidate phosphohydrolase</td>
<td>173</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>174</td>
</tr>
<tr>
<td>Diacylglycerol acyltransferase (Diacylglycerol O-acyltransferase; diglyceride acyltransferase)</td>
<td>DAGAT</td>
<td>2.3.1.20</td>
<td>Acyl-CoA:1,2-diacylglycerol O-acyltransferase</td>
<td>175</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>176</td>
</tr>
<tr>
<td>Choline phosphotransferase (Diacylglycerol cholinephosphotransferase; phosphorylcholine–glyceride transferase)</td>
<td>CPT</td>
<td>2.7.8.2</td>
<td>CDPcholine:1,2-diacylglycerol-cholinephosphotransferase</td>
<td>177</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>178</td>
</tr>
<tr>
<td>Lyso phosphatidylcholine acyltransferase (1-Acylglycerophosphocholine O-acyltransferase; lyssolecithin acyltransferase)</td>
<td>LPCAT</td>
<td>2.3.1.23</td>
<td>Acyl-CoA:1-acyl-sn-glycerol-3-phosphocholine O-acyltransferase</td>
<td>179</td>
</tr>
</tbody>
</table>

$^a$EC numbers from Ref. 37.

*Source*  Ref. 37 and other works indicated.
Figure 5  Triacylglycerol assembly reactions occurring in plant endoplasmic reticulum. Details are the same as in Figure 4.
of reactivity is also supported by an *Arabidopsis* mutant lacking *fad2* (165). The reactivity of 18:2\((\Delta 9,12)\)-PC \(\Delta 15\) desaturase (\(\Delta 15\)DES) is almost identical to that of 18:2\((\Delta 9,12)\)-PC as just described for the \(\Delta 12\)DES on 18:1\((\Delta 9)\)-PC (166). However, there is also evidence for multiple forms of the \(\Delta 15\)DES, one with reactivity toward both positions and the other reactive only on the sn-2 18:2\((\Delta 9,12)\) residue, leading to an enrichment in 18:3\((\Delta 9,12,15)\) at the sn-2 position of glycerolipids. The \(\Delta 12\)DES and \(\Delta 15\)DES are not capable of recognizing 18:2\((\Delta 6,9)\) and 18:3\((\Delta 6,9,12)\), respectively, as substrates (14).

A less common membrane desaturase occurs in seeds of borage (*Borago officinalis*), and also likely, evening primrose (*Oenothera biennis*) and black currant (*Ribes nigrum*), all of which accumulate \(\gamma\)-linolenic acid (18:3\((\Delta 6,9,12)\)) in seed oils (Table 2). The \(\Delta 6\) double bond is introduced by a \(\Delta 6\)-desaturase (\(\Delta 6\)DES) located in the endoplasmic reticulum (Fig. 4, Refs. 167 and 168). The \(\Delta 6\)DES also requires \(O_2\) and NADH as cosubstrates and acts on PC-linked 18:2\((\Delta 9,12)\). However, in contrast to the positional selectivity of the \(\Delta 12\)DES and \(\Delta 15\)DES, the \(\Delta 6\)DES from borage recognizes only the sn-2 residue of 18:2\((\Delta 9,12)\)-PC for the desaturation reaction.

The role of the desaturases in triacylglycerol assembly is elucidated further in Sec. V. Despite inability to purify these desaturases, the identification of *Arabidopsis* mutants (15,16) has allowed genetic transformation of various hosts to proceed in attempts to modify the profile of unsaturated fatty acids in seed/fruit oils (see Sec. VI).

**V. TRIACYLGLYCEROL ASSEMBLY**

**A. Overview**

Following plastidic biosynthesis (Fig. 1), and modification of the acyl chains in the plastid (Fig. 3) and endoplasmic reticulum (Fig. 4), the various fatty acids are finally destined for incorporation into storage (triacylglycerols) or functional (polar) glycerolipids. In oil-bearing seed or fruit tissues, assembly of lipids of both types takes place in the endoplasmic reticulum (some polar glycerolipid assembly also takes place in the plastid of “16:3” plants (35) (Sec. III B.7 and Fig. 3). With both functional and storage lipids assembled in the endoplasmic reticulum, it quickly becomes obvious that mechanisms must exist to allow for discrimination in the pool of fatty acids to be used for these divergent purposes. Despite the diversity of fatty acids in the plant kingdom (Table 2), a strict fatty acid profile among the polar glycerolipids that are destined to be functional unit in cellular membranes must be maintained to ensure survival of the organism (10,13,14). Thus, the divergent pathways of storage and functional lipid assembly must have safeguards to prevent nonproprietary utilization of fatty acids.

There are two basic routes of triacylglycerol synthesis or assembly (Fig. 5). One route involves the exclusive use of the so-called Kennedy (169) pathway, which operates in a rather linear fashion, drawing from the acyl-CoA pool to satisfy the needs for triacylglycerol assembly. The other route for triacylglycerol assembly uses the Kennedy pathway as modulated by the involvement of phosphatidylcholine (PC) and its derivatives. Different organisms evoke these pathways to different degrees, and this is reflected in their characteristic triacylglycerol compositions. This section first describes regulatory features of triacylglycerol assembly and then examines the
characteristic processes in seeds or fruit of plant species that collectively accumulate triacylglycerols composed of (a) unsaturated (Δ9,Δ12,Δ15) 18-carbon fatty acids, (b) medium chain (<16 acyl carbons) fatty acids, or (c) long chain (>18 acyl carbons) or otherwise unusual fatty acids. It is hoped that the reader will emerge with a clear picture of the nature of the nonrandom distribution of fatty acids in plant storage triacylglycerols.

B. Control of Triacylglycerol Assembly

Regulation of triacylglycerol assembly is vested in the individual enzymes involved (see Table 5 and Fig. 5). The Kennedy pathway comprises,* in sequence, GPAT (170), LPAAT (171,172), PTAP (173,174), and DAGAT (175,176). These steps sequentially mediate the addition of acyl groups from the acyl-CoA pool to the glycerol backbone. The involvement of PC-linked acyl groups involves principally CPT (177,178) and LPCAT (179) [or in some cases, phospholipase A (147)]. The latter three enzymes provide access of 18:19 from the CoA pool, via PC, to Δ12DES and Δ15DES activities to yield PC-linked 18:29,12 and 18:39,12,15. These polyunsaturated fatty acids can be used in triacylglycerol assembly either via return to the acyl-CoA pool (by LPCAT or phospholipase A) or PC-diacylglycerol exchange (CPT) followed by DAGAT action. PC is originally assembled by the Kennedy pathway involving GPAT, LPAAT, PTAP, and CPT (15). [The author has not located any reports of discrete isoforms or pathways exclusive for diacylglycerol or PC assembly, although the presence of divergent pathways has been implied (20)]. In any event, the salient features of involvement of PC in triacylglycerol assembly revolve around the role of PC in mediating the desaturation and oxygenation modifying steps.

The features that control channeling of fatty acids toward triacylglycerol assembly are (a) the level and specificities of the enzymes mediating each acylation step, (b) the nature of the acyl-CoA pool as imported from the plastid or modified within the endoplasmic reticulum, (c) the exchange of PC-linked fatty acids with the acyl-CoA pool, and (d) the extent to which exchange takes place between the PC and diacylglycerol pools. Features a and b are interdependent such that for plant species where the Kennedy pathway enzymes are relatively nonselective, the profile of the acyl-CoA pool dictates the types of the fatty acyl residue evolving as components of triacylglycerols. On the other hand, for species where Kennedy pathway enzymes exhibit pronounced selectivity, the acyl-CoA pool profile has less influence on which fatty acyl groups evolve as components of triacylglycerols. When Kennedy pathway enzyme selectivity and the acyl-CoA pool are matched, up to 90% of the fatty acyl groups evolving as components of triacylglycerols can be a single species.

Less attention has been paid to which enzyme(s) regulate triacylglycerol assembly than to a characterization of the enzyme reactions composing each step. Based on a comparison of relative magnitudes of in vitro activities in microsomal membranes of safflower (14), the most limiting enzyme activities are DAGAT, CPT (PC ⇒ diacylglycerol direction), and Δ12DES, and the next most limiting are CPT (diacylglycerol ⇒ PC direction), PTAP, and GPAT. Extrapolating from animal sys-

*See Table 5 for abbreviations.
tems, PTAP is a likely candidate for regulatory control (35), and the 18:1\textsubscript{\omega-9}-induced translocation of PTAP from the cytosol to the endoplasmic reticulum of maturing safflower seeds was suggested to be a feedward control mechanism (173,174). DAGAT has also been suggested as being rate-limiting in rapeseed (35) and is a candidate enzyme for overall control of flux toward triacylglycerol synthesis (14), since it is the only enzymic step dedicated to triacylglycerol assembly (109). In species where storage oils are dominated by 18:1\textsubscript{\omega-9}/18:2\textsubscript{\omega-9,12}/18:3\textsubscript{\omega-9,12,15}, CPT is suggested to control flux of fatty acids and glycerol toward triacylglycerol assembly (177,178). The acyl-CoA pool may also have regulatory features, since it induces import of PTAP (174), and acyl-CoA levels and fatty acid profile have impact on rates of some enzyme activities in the Kennedy pathway (180). Finally, two metabolically discrete pools of diacylglycerol are suggested to exist (181), potentially exerting control over flux toward triacylglycerol synthesis. In any case, triacylglycerol accumulation is a developmentally regulated process that is synchronized with the maturation of seed or fruit tissue (35,148) and must involve coordinated expression of seed/fruit-specific genes.

C. Seed Oil Enriched in Unsaturated (\Delta9, \Delta12, \Delta15) 18-Carbon Fatty Acids

1. General Aspects

In these species [soybean, safflower, sunflower (Helianthus annuus), avocado, cocoa (Theobroma cacao), maize (Zea mays), palm, and linseed (Linum usitatissimum)], triacylglycerol assembly results in the occupation of the sn-1,3 positions by 16:0/18:X, and of the sn-2 position by 18:X. Arabidopsis is a good genetic model for these species because it accumulates 30\% 18:2\textsubscript{\omega-9,12}, 20\% 18:3\textsubscript{\omega-9,12,15} (in addition to 22\% 20:1\textsubscript{\omega-11}) in seed oil, and the genetics of this organism are well known (182). Species accumulating unsaturated fatty acids with double bonds in positions other than \Delta9, \Delta12, and \Delta15, are discussed as a group in Sec. V.E.

2. Involvement of Kennedy Pathway

Generally, GPAT has broad specificity, as characterized by the safflower enzyme (170,183), although it best recognizes 16:0/18:X-CoA substrates. The GPAT from cocoa beans is slightly different in that only saturated 16/18-CoA substrates are recognized and 18:1-CoA is not very reactive (184). Because GPAT is relatively nonselective, the fatty acyl residue appearing at the sn-1 site will largely be representative of the acyl-CoA pool available, which in turn is a reflection of the primary products of FAS and modification reactions.

In contrast, LPAAT, is characterized from cocoa bean, palm endosperm (kernel), maize, rape, and safflower, tends to be very selective for unsaturated 18:X-CoA substrates (171,176,183,184). Thus, the combined selectivities of GPAT and LPAAT confer the “eukaryotic” configuration of glycerolipids in these species. Furthermore, LPAAT is selective also for the species of lysophosphatic acid (LPA) cosubstrate in that maize and rape LPAAT prefer 18:1\textsubscript{\omega-9}-LPA as substrate and will react with 12:0-LPA (176). On the other hand, palm endosperm LPAAT will recognize both 12:0- and 18:1\textsubscript{\omega-9}-LPA, but will tend to couple 12:0-LPA with another 12:0 residue, whereas no selectivity between 12:0- and 18:1\textsubscript{\omega-9}-CoA is observed with 18:1\textsubscript{\omega-9}-LPA as substrate (176). This property helps partition unusual or “nonfunctional” fatty
acids in a manner that induces them to evolve into triacylglycerols (see Sec. V.D for other aspects of medium chain triacylglycerol assembly).

The next step in the Kennedy pathway, PTAP, has not been well studied, but in safflower it shows greater selectivity for phosphatidic acid (PA) with two unsaturated acyl groups than for PA species with one or no unsaturated acyl groups (185). This pattern of selectivity probably affords little influence on triacylglycerol composition, but it may help ensure that saturated fatty acids are not allowed to readily gain entrance into the PC pool via subsequent CPT action.

Finally, DAGAT from maize, castor, and groundnut, and probably many other oilseed species, like GPAT, exhibits rather broad specificity among 14:0/16:0/18:0/18:X-CoA substrates (175,176). Thus, the fatty acyl groups evolving at the sn-3 position are largely defined by the profile of the available acyl-CoA pool. Palm mesocarp is unusual in that the DAGAT associated with the developing oil bodies (derived from the endoplasmic reticulum, see Sec. V.F regarding oil body genesis) is selective for 18:1_{3\alpha} over 16:0-CoA compared to the broad specificity of endoplasmic reticular DAGAT activity (176). This implies the presence of two populations of DAGAT (perhaps conferred by the membrane environment), with that in developing oil bodies selecting for unsaturated 18:X acyl species at sn-3 for triacylglycerols being actively deposited in accumulating oil bodies. Some sources of DAGAT have more pronounced selectivities; that of cocoa beans, for example, is selective for 16:0-CoA and 18:0-CoA substrates (184), contributing to the characteristic triacylglycerol species in this species, which is 24–30% 16:0 and 30–36% 18:0 (28). The unusual enrichment of 18:0 in storage triacylglycerols in cocoa beans must be accounted for, in part, by the accumulation of 18:0-CoA in the acyl-CoA pool. Thus, during development or maturation of cocoa beans, some mechanism is evoked whereby Δ9DES activity is suppressed (and/or TE action toward 18:0-ACP is enhanced), allowing 18:0 to be produced at levels much beyond the normal few percent that typically occurs in plant tissues. This yet-to-be-identified trait may constitute an alternative approach to manipulating plant species to produce high 18:0 oils, which in turn could circumvent the need for hydrogenation as a means to yield margarine-type fats (see Sec. VI.B).

3. Involvement of PC Exchange Reactions

Perhaps the most important enzyme activities in these plant species are CPT and LPCAT, which mediate steps in triacylglycerol assembly that allow 18:2_{3\alpha,1\beta}/18:3_{3\alpha,1\beta,15} to accumulate (177,181,186). In concert with the Kennedy pathway, and when 18:1_{3\alpha} is in abundance (as would be expected from many plant FAS systems), the collection of these activities gives rise to triacylglycerols with mono- and polyunsaturated 18:X fatty acyl groups occupying all three sites of the triacylglycerols (instead of 16:0/18:0 residing at one or both of the primary sites). In both sunflower and safflower seeds, 18:1_{3\alpha} is esterified at sn-1 and sn-2 positions by GPAT and LPAAT, respectively. Following LPAAT action, which also favors esterification of 18:1_{3\alpha}, the resulting diacylglycerol can exchange with the PC pool via CPT. The PC-linked 18:1_{3\alpha} is now subject to desaturation by Δ12DES and Δ15DES, which are relatively nonspecific for position (sn-1 or -2) of the acyl group (178). The resulting PC derivatives can undergo a reciprocal exchange (via CPT) to form diacylglycerol, which is then subject to DAGAT action to yield the final triacylglycerol product. Alternatively, the presence of LPCAT, which is highly selective for ester-
ifying 18:1<sub>9</sub> and deacylating 18:2<sub>9,12</sub>/18:3<sub>9,12,15</sub>, allows for reversible exchange between the acyl-CoA pool and PC-linked acyl groups (179). Thus, after desaturation, 18:1<sub>9,12</sub> and 18:3<sub>9,12,15</sub> may be released to the acyl-CoA pool to be utilized in other assembly steps of the Kennedy pathway recognizing 18:2<sub>9,12</sub>- and 18:3<sub>9,12,15</sub>-CoA as substrates. In safflower, sunflower, and linseed, these fatty acids appear to be utilized by GPAT and LPAAT (187). Perhaps, more importantly, 18:2<sub>9,12</sub>- and 18:3<sub>9,12,15</sub>-CoA can be used by DAGAT to complete the acylation of triacylglycerols exclusively with (poly)unsaturated 18:X residues, a trait characteristic of seed oils of many of these species (28). For example, linseed oil has 50–60% of each acyl site occupied by 18:3<sub>9,12,15</sub>.

4. Unique Features of Species in This Group

In keeping with the triacylglycerol profile of cocoa butter and avocado oil, the activities of LPCAT and CPT are not very involved in triacylglycerol assembly: there is little 18:2<sub>9,12</sub> in these oils, and it occurs primarily in the sn-2 position (28). It should be noted that diacylglycerol and PC formation share a common (Kennedy) pathway, and PC with polyunsaturated 18:X residues is required for assembly of functional (membrane) lipids in these plant species. Since it is so vital for the plant cell to segregate fatty acids into storage and functional lipid, it is somewhat perplexing that CPT (such as in rape and sunflower) exhibits little selectivity (181) and therefore cannot serve as a “gatekeeper” to exclude unusual fatty acids from the PC pool. In view of the combined broad specificity of DAGAT, it has been argued that two pools of diacylglycerol exist, and these are spatially (or metabolically) segregated to satisfy the divergent needs of storage and functional lipid assembly (181). Considering the complexity of all the acyl modification and glycerolipid assembly reactions that take place in the endoplasmic reticulum, it would be surprising if no metabolic compartmentation existed for some of these processes.

Finally, the importance of the composition of the acyl-CoA pool, as a contributor to the resulting profile of assembled triacylglycerols, cannot be overstressed. Even though safflower seed oil is about 90% 18:1<sub>9</sub> and 18:2<sub>9,12</sub> (28), when presented with equimolar levels of 16:0/18:0/18:1<sub>9</sub> in vitro, the (non)selectivity features of the enzymes involved in the Kennedy pathway can give rise to cocoa butter–type fats (notably rich in 16:0 and 18:0) (183). This is the type of biosynthetic potential that presents opportunities for exploitation in manipulating plant species for specific end uses in food.

D. Oils Enriched in Medium Chain Fatty Acids

1. General Aspects

Species important in this category are coconut, palm (kernel), and the genus Cuphea (27,28). In some of these species, more than 70% of the acyl groups in seed oil can be a single fatty acid, indicating that these “unusual” fatty acids occupy all three sites of the triacylglycerol. Perhaps the most comprehensive study on triacylglycerol assembly in this group of oil-bearing species was derived for Cuphea lanceolata (188), which accumulates up to 83% 10:0 (27). It should be remembered that in these species, the existence of TE isoforms allows premature termination of FAS and export of these fatty acids (acyl chain lengths of <16) to the endoplasmic reticulum (see Sec. III.B.7). Thus, the acyl-CoA in these species can be expected to be
very enriched in medium chain length fatty acids, especially during seed development/maturation.

2. Involvement of Kennedy Pathway

The GPAT activities in *C. lanceolata*, *C. procumbrens*, and *C. wrightii* are characterized by selectivity toward 10:0–14:0-CoA > 18:X/16:0-CoA substrates (188,189). Perhaps the most critical step in these species is the LPAAT reaction. For 12:0/14:0 LPA species, acyl-CoA of fewer than 16 carbons are utilized as substrates for the second esterification step, resulting in PA products with medium chain fatty acyl residues at both sn-1 and sn-2 sites, and this pattern also occurs for palm endosperm LPAAT (166). When 18:1ω9-LPA is available for reaction, LPAAT exhibits a strict preference for another 18:1ω9-CoA to become esterified to form di-18:1ω9-PA in *Cuphea* spp. (188,189), or in the case of palm kernel LPAAT, either 12:0- or 18:1ω9-CoA is used to yield di-18:1ω9-PA and 12:0,18:1ω9-PA (172). In coconut endosperm, 10:0/12:0/14:0-CoA is preferred for reactions with both 12:0/18:1ω9-LPA species (190,191). In palm endosperm, 18:1ω9-LPA is preferred to 12:0-LPA (172), whereas in coconut endosperm the reverse is the case (190).

The result of LPAAT and subsequent PTP action in *Cuphea* spp. is a bimodal distribution of di-18:1ω9-acylglycerol and di-10:0/12:0/14:0-acylglycerol (188). DAGAT is selective for the latter diacylglycerol species and specifically utilizes another 10:0/12:0/14:0-CoA for the final step in assembling triacylglycerol (181,188). The di-18:1ω9-acylglycerol is almost exclusively used for exchange (CPT) to the PC pool for incorporation into functional lipids. Thus, the combined selectivities of the GPAT, LPAAT, and DAGAT strictly segregate fatty acids into storage and functional glycerolipids (188,192). The CPT enzyme is not believed to play a major role in excluding various diacylglycerol species from the PC pool (181). Rather, the lack of di-10:0/12:0/14:0-PC in the PC pool may be accounted for in the comparatively high DAGAT activity toward di-10:0/12:0/14:0-acylglycerol (and rapid channeling into triacylglycerol), or the existence of two discrete metabolic pools of diacylglycerol: one for PC exchange and the other for triacylglycerol assembly (181).

E. Oils Enriched in Unusual Fatty Acids

1. General Aspects

Very long chain fatty acids (VLCFAs), oxygenated fatty acids [such as ricinoleic acid (18:1ω12-OH), and vernolic acid (18:1ω9,12/13-epoxy)], and those with unusual double-bond positions (Δ5, Δ6, Δ11) are treated as a group. An understanding of how these unusual fatty acids are channeled in the host plants is useful in building a comprehensive view of triacylglycerol assembly in all plants and provides insight into genetic traits that may be worthwhile to exploit in plant oils used for foods.

VLCFAs include those of 20–24 acyl carbons. The most notable are erucic (22:1ω13) and gondoic (20:1ω11) acids in rapeseed, and the same fatty acids plus 20:1ω5, 22:2ω5,ω13 in meadowfoam, where the Δ5 double bond arises from Δ5DES action after ACE-mediated elongation from the 16:0/18:1ω9 parent fatty acid (193). Other species relevant to this grouping include mustard (*Sinapis alba* and *Brassica* spp.), *Crambe* spp., groundnuts (source of 20:0, 22:0, and 24:0), honesty (*Lunaria annua*, source of nervonic acid, 24:1ω5), borage, black currant and evening primrose.
oils (rich in 18:3_6,9,12), and coriander (and other Umbelliferae, rich in 18:1_6,9,12) (24–26,28); Δ6DES and Δ4DES activities account for the presence of these latter two groupings of species. These and other structurally related fatty acids are strictly excluded from functional glycerolipids, and in the case of seed oils from rape, mustard, *Crambe* spp., groundnuts, and honesty, the VLCFAs are virtually confined to the sn-1,3 positions in triacylglycerols (28,194); this restricts the present capability of trying to develop for industrial purposes high-erucic rape exceeding the theoretical 67% yield of erucic acid (145,146,148).

2. Involvement of Kennedy Pathway

By all accounts the Kennedy pathway is utilized for triacylglycerol assembly in these species (194,195). *GPAT* has broad selectivity toward acyl-CoA substrates (180,196) and will readily act on 22:1_13-CoA but shows no preference for it. Thus, the profile of the acyl-CoA pool largely dictates which acyl group is attached to the sn-1 position (180). As expected, *LPAAT* is selective toward unsaturated 18:X-CoA substrates and is responsible for the exclusion of 22:1_13 (and like fatty acids) from the sn-2 position in the transient PA pool in rape (180,196,197). This same specificity applies to palm, castor, soy, and maize *LPAAT* (197). Although the *GPAT* and *LPAAT* of castor bean have not been studied in detail, it could be hypothesized, based on other sources of these enzymes, that they are rather broad in selectivity and that the tendency for castor bean to yield triricinolein is controlled to a large degree by the abundance of 18:1_9,12-OH in the acyl-CoA pool.

**DAGAT** has pronounced selectivity for 22:1_13-CoA as substrate [even in low-erucic rapeseed oil, since it is the elongase that is dysfunctional and not the **DAGAT** (35,180)], but it also prefers to react with di-18:1_19-, or di-18:2_9,12-acylglycerols over mixed 18:X/22:1_13-diacylglycerols (181). Again, the acyl-CoA pool would apparently have substantial influence on the acyl-CoA and diacylglycerol species available for reaction with the final **DAGAT** step. **DAGAT** from borage was inferred to be highly selective for 18:3_6,9,12 appears at sn-1, whereas it comprises 50% sn-3 acyl groups in mature seed oil; this composition helps to ensure that 18:3_6,9,12 is dedicated to triacylglycerol and not functional lipid assembly (168). **DAGAT** of castor bean is highly selective for di-18:1_9,12-OH-acylglycerol over other 18:X-diacylglycerol species and completes the assembly of triacylglycerols rich in 18:1_9,12-OH (147).

Flux of acyl-CoA and glycerol through the Kennedy pathway toward triacylglycerol assembly in these plant species is rapid, inasmuch as intermediates with 22:1_13 residues do not appreciably accumulate but are rapidly labeled with [14C]acyl-CoA substrates (194,195). The PC pool is also involved in triacylglycerol assembly, likely where 18:2_9,12/18:3_9,12,15 residues appear at the sn-2 position of oils from these species.

3. Involvement of PC Exchange Reactions

Utilization of oxygenated fatty acids in triacylglycerol assembly has been best studied in the beans of castor, which accumulate over 90% as ricinoleic acid. The utilization (indeed, genesis) of oxygenated fatty acids requires access of the parent fatty acid to the PC pool. The **12-MO** that forms ricinoleic acid in castor bean (147), and the **12-EPOX** system to form vernolic acid (18:1_9,12,15-epoxy), which accumulates to levels of 70% in *Euphorbia lagascae* (149), require PC-linked 18:1_9, and specifically, sn-
2-18:1<sub>6</sub>-PC for the 12-MO. Upon hydroxylation of the 18:1<sub>6</sub>-PC in castor bean, the ricinoleic acid is released quickly by phospholipase A<sub>2</sub> to form 18:1<sub>6,12</sub>-OH (and ultimately its -CoA derivative) and lyso phosphatidylecholine (LPC) (147). LPC is reacylated by LPCAT, which is selective for 18:1<sub>6</sub>-CoA. The repetition of this cycle allows for ricinoleic acid to accumulate in the acyl-CoA pool and exclude it from the PC pool and functional lipids. Ricinoleoyl-CoA is formed and can be used successively in GPAT, LPAAT, and DAGAT reactions to yield the tri-18:1<sub>6,12</sub>-acylglycerol.

4. Unique Features of Species in This Group

A notable exception of the trends of triacylglycerol assembly in these plant species rests with the LPAAT of meadowfoam. The meadowfoam LPAAT can utilize 22:1<sub>13</sub> and is most reactive with mono-22:1<sub>13</sub>-acylglycerol (196,197). The potential to utilize this trait in transgenic rape to favor the formation of trierucin is an opportunity currently being evaluated (198,199).

This process by which unusual fatty acids are channeled into storage lipid in coriander and carrot endosperm (and related species that can accumulate up to 85% of storage triacylglycerol as 18:1<sub>6</sub>; (see Table 2, Refs. 27 and 28), appears to be different from that for other species in this group. In carrot and coriander endosperm, the PC-diacylglycerol exchange pathway (CPT step) constitutes a major route for triacylglycerol assembly, since newly synthesized (and radiolabeled) fatty acids (including 18:1<sub>6</sub>) become most immediately concentrated in PC, and then in triacylglycerol (200). This observation implies a role for PC in triacylglycerol assembly beyond its well-documented role of simply allowing for acyl modification reactions to occur.

The general enzymic patterns of triacylglycerol assembly in species belonging to the groups discussed above (Secs. V.C–V.E) is offered as a summary in Table 6.

F. Oil Body Genesis

Some of the earliest in vitro experiments demonstrating triacylglycerol accumulation mediated by safflower microsomal membranes yielded a milky-white suspension (201). Upon centrifugation of the reaction mixture, a buoyant “fat pad” was formed at the surface, and electron microscopic examination of this material indicated the presence of “naked” fat globules. These oil bodies or “oleosomes” contained detectable activities of some of the enzymes involved in triacylglycerol assembly (202).

Since these initial studies, a reasonable view of genesis of oil bodies in plant seed (and pollen) tissues has evolved (145,146,203,204). Oil bodies (~20 μm diameter) form as vesicles that “bud off” from the endoplasmic reticulum, and this process may be induced by triacylglycerol accumulation and the attendant swelling or distension of the membrane. This is not a universal view of oil droplet formation, for there is evidence of a cytoplasmic origin for seed/endosperm tissues of some species. However, these apparently discordant views collectively provide the basis for a growing realization that there may be distinct subpopulations of the endoplasmic reticulum, each with a different profile of enzyme activities with lipid metabolism. This view is supported by the body of studies in which microsomal fractions have been prepared from various tissues at varying degrees of maturation, with findings of varying levels and profiles of marker enzymes. This view of endoplasmic
reticulum subpopulations also finds support in the observation that different DAGAT reaction selectivities exist in oil bodies and endoplasmic reticulum preparations from palm mesocarp (176), and in the suggestion that discrete metabolic pools exist for diacylglycerol intermediates in the divergent pathways for storage and functional lipid assembly (181).

Immature (newly formed) oil droplets are less than 0.2% protein, and they can coalesce because they are only partially stabilized by a phospholipid/protein annulus (145,146,203). Stable oil bodies are formed upon deposition of seed-specific proteins, “oleosins” (205,206), which can account for up to 20% of total seed protein during seed development (207). These oleosins, which are tightly bound to oil droplets, are believed to function principally to stabilize the oil droplets, and not to be involved in their original formation (145,146,203). The mechanism of oleosin deposition is believed to involve an enrobing of naked oil droplet by the endoplasmic reticulum.

Oleosins are small (~19 kDa) and characterized by three domains roughly equal in size: a C-terminal polar region, a central hydrophobic region, and an N-terminal amphiphilic region. The terminal domains are oriented toward the cytoplasmic face, and the hydrophobic domain, which is highly conserved among species and is of β-strand character, anchors the protein firmly into the oil droplet (145,146,203). Oleosins can be classified by homology as belonging to the pollen or seed groups, and there is an apparent lack of oleosins in oil bearing fruits such as avocado, coconut, olive, and palm, for reasons unknown at this time.

An eventual application of the molecular biology of these oleosins will likely be the preparation of fusion proteins or oleosin adducts that contain valuable peptides (148,204). These valuable peptides could be liberated and recovered by subsequent hydrolysis of the peptide linkage after recovery of the adducts by simple flotation of oil bodies following disruption of the host tissue. The ease of recovery of oil droplets and the better than 85% targeting of the oleosin–peptide adduct to the oil droplets, even for large (65 kDa) peptides, make this a promising and economical approach to large-scale production of biochemicals. This approach also could be used as an alternative enzyme immobilization technique, where the enzyme remains anchored to the oil droplet by virtue of its covalent linkage to the oleosin (208).

VI. GENETIC APPLICATIONS AND OPPORTUNITIES FOR LIPID MODIFICATION

A. Current Status and Overview

As of this writing, genes have been cloned, from various sources, for almost all the proteins (viz., oleosins, ACP) and enzymes involved in fatty acid biosynthesis and modification, glycerolipid assembly, and oil body genesis (20,93,109,209–211). It is reasonable to expect that the genes that have not yet been cloned will be cloned within the next few years. Thus, we can soon expect to have the complete spectrum of capability for genetically manipulating plant species for any exploitable purpose. However, a dilemma exists in that the rapid progress made in understanding plant storage (and functional) lipid biosynthesis by genetic approaches in the last decade has outpaced the rate at which advances are made in understanding how lipid structure–function impacts, contributes to, or can be used to otherwise control, food quality. In short, food scientists (and nutritionists) need to identify the problems and
**Table 6** Summary of Trends of Enzyme Selectivity and/or Involvement in Triacylglycerol Assembly in Oil-Bearing Plant Tissues Rich in Various Groups of Fatty Acids

<table>
<thead>
<tr>
<th>Enzymic step (control point or feature)</th>
<th>Oil-bearing plant species/tissues accumulating triacylglycerols rich in</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Unsaturated 18-carbon fatty acids</td>
</tr>
<tr>
<td>GPAT (sn-1-acyl)</td>
<td>Broad specificity; selective toward 16:0/18:X-CoA</td>
</tr>
<tr>
<td>LPAAT (sn-2-acyl)</td>
<td>Selective toward unsaturated 18:X-CoA and 18:X-LPA</td>
</tr>
<tr>
<td>PTAP</td>
<td>Selective toward diunsaturated-PA</td>
</tr>
<tr>
<td>DAGAT (sn-3-acyl)</td>
<td>Broad specificity&lt;sup&gt;3&lt;/sup&gt;</td>
</tr>
<tr>
<td>Table Entry</td>
<td>Extensive involvement,\textsuperscript{a} Not selective; DAGAT selectivity leaves di-18:1\textsubscript{α,δ}-acylglycerol for exchange into PC</td>
</tr>
<tr>
<td>-------------</td>
<td>---------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>CPT (DAG ↔ PC)</td>
<td>Extensive involvement.\textsuperscript{a} Not selective; DAGAT selectivity leaves di-18:1\textsubscript{α,δ}-acylglycerol for exchange into PC</td>
</tr>
<tr>
<td>LPCAT (PC ↔ lysoPC) (or PLA\textsubscript{2})</td>
<td>Extensive involvement; selective toward 18:1\textsubscript{α,δ}-CoA for acylation; selective toward 18:2\textsubscript{α,β,γ}/18:3\textsubscript{α,β,γ,δ} for deacylation; contributes 18:X to acyl-CoA pool</td>
</tr>
<tr>
<td>Acyl-CoA pool</td>
<td>Important for GPAT and DAGAT steps</td>
</tr>
</tbody>
</table>

\textit{Notes} (comments or notable exceptions)
\textsuperscript{a}Cocoa bean enzyme is selective for 16:0/18:0-CoA.
\textsuperscript{b}Palm kernel enzyme recognizes 12:0-LPA.
\textsuperscript{c}Cocoa bean/borage enzymes have greater selectivity.
\textsuperscript{d}Except in cocoa bean and avocado.
\textsuperscript{e}Palm kernel and coconut enzymes also form mixed 12:0/18:1\textsubscript{α,δ}-PA species as major products.
\textsuperscript{f}Meadowfoam enzyme is an exception.
\textsuperscript{g}Inferred from properties of GPAT from plant tissues in general.
\textsuperscript{h}Inferred from properties of LPAAT from plant tissues in general.
\textsuperscript{1}Abbreviations for enzymes and fatty acids are the same as in Table 5 and footnote 1 (Sec. II); respectively. Other abbreviations: PA, phosphatidic acid; LPA, lysophosphatidic acid; DAG, diacylglycerol; PC, phosphatidylcholine; LPC, lysophosphatidylcholine; PLA\textsubscript{2}, phospholipase A\textsubscript{2}, N/A, not appropriate or not available.
limitations associated with using native lipids in foods on a molecular level, if the power of molecular biology is to be harnessed to furnish solutions to these problems.

Chapter 30 presents a discussion of the current status of genetic engineering and its application to food lipid modification. There are also many examples of transgenic strategies for lipid modification for industrial purposes (111,148,210,212–214). It is hoped that the preceding pages in this chapter have placed the reader in a position to clearly understand the strategy or basis for the genetic approaches discussed in this section and in Chapter 30.

This section introduces current applications and opportunities that exist for the genetic manipulation of plant lipids for specific food uses. The examples are also intended to illustrate the need for food scientists to be able to identify the specific features of lipid structure–function relationships that have impact on food quality attributes. Such applications can be delineated as opportunities to improve food lipids, principally triacylglycerols, from a physical, chemical, or nutritional perspective.

B. Modifying Physical Functionality

Opportunities to improve physical properties of bulk lipids rest with modification of the acyl chain profile and positional distribution within triacylglycerols (33). Current activities in this area (215) include studies of transformed plants that yield seed oils enhanced in saturated fatty acids (either 12:0 or 18:0).

Enhancing 18:0 content in storage triacylglycerols provides an alternative to chemical hydrogenation (and the attendant and persistent, but nebulous, threat of the presence of trans fatty acids) in the preparation of margarines and table spreads (210,211,216). The basis for enhancing 18:0 levels is antisense technology directed at the gene encoding for Δ9DES, as applied in soybean and canola (217,218). This results in greater steady state levels of 18:0-ACP accumulating and the enhanced opportunity for TE to act on 18:0-ACP, as well as release of 18:0 for export to the endoplasmic reticulum for the purpose of triacylglycerol assembly (cf. Figs. 3 and 5). High-18:0 oils from soybean transformants have been evaluated for use in margarine products, and while admixture with other solid fats is necessary to obtain appropriate physical properties, the use of the oils from transgenic soybeans can be considered to be at least partially successful in this regard (219).

Another genetic approach to achieving this outcome is to delete the gene encoding for Δ9DES, and this has been done for Arabidopsis thaliana (220,221). Although there was an increase in 18:0 in the seed oil of the mutants, there was also an increase in 18:0 of leaf lipids, and this led to aberrant morphology, representing a negative impact on plant development. This set of results illustrates the need to specifically target seed-specific genes, so as to not impair any agronomic traits (148) or incur any “agronomic penalty” (222).

One approach that has not been exploited is to elucidate, and then genetically manipulate, the factors that exert developmental control over lipid synthesis in plants oils known to accumulate 18:0 (cf. Secs. V.C.2 and V.C.4), such as cocoa bean (28,184,223). Such an understanding could potentially identify alternative strategies for enhancing saturated fatty acid content of storage triacylglycerols in other species. Recall that safflower seed enzymes involved in triacylglycerol assembly also possess selectivities that in vitro can give rise to the dominant triacylglycerol molecular species in cocoa butter, provided the appropriate profile of acyl-CoA substrates is
available (cf. Sec. V.C.4 and Ref. 183). Any of these genetic strategies would augment the more conventional (and reactionary!) approach of preparing cocoa butter substitutes from various lipid sources via postproduction enzymic modification using lipases (see Chapters 26 and 27). Second, from a mechanistic viewpoint, understanding the influence of temperature on triacylglycerol assembly in cocoa beans may allow for identification of a means to ameliorate the problem of “soft” cocoa butters that originate in warmer climates (223).

With regard to other biotechnological approaches to producing cocoa butter, plant cell (suspension) cultures have been investigated for this purpose (224). Unfortunately, nondifferentiated cells produce lipids that resemble functional (and not storage) lipids in fatty acyl composition. This can be easily explained in terms of developmental and organ-specific regulation of storage lipid accumulation. Tissue differentiation during culture was previously shown to be important in that maturing cacao embryos grown in vitro did accumulate limited amounts of triacylglycerol with a molecular species profile resembling that of cocoa butter (225).

A lot of interest has been devoted to preparing and testing transgenic plants that produce 12:0-rich oils, specifically in canola (215,226). Such oils would serve as replacements for selected tropical oils (viz., coconut, palm kernel) and could be used in applications such as cocoa butter extenders, coatings, simulated dairy products, bakery goods, and spreads. The apparent excitement about the development of a canola strain that produces lauric acid is somewhat perplexing, considering the campaign of “tropical-oil bashing” (also including palm oil) in North America that was initiated in the late 1980s (227,228). Based on the contribution of tropical oils to the diet of the U.S. public, the slight hypercholesterolemic effect of 12:0 (22°) is probably dismissible (230), and in the case of palm oil, not at all relevant (231) (because palm oil essentially lacks 12:0). However, the negative image of 12:0-rich oils is still apparent, since the need to address this issue with 12:0-producing transgenic canola was met with a comment suggesting that the canola transformant stores 12:0 in a “novel form” (2320. This speculation was apparently based on the difference in LPATT activities of rapeseed and coconut, in that the former excludes, while the latter permits, the positioning of 12:0 at the sn-2 site (see Secs. V.D.2 and V.E.2, Table 6, and Ref. 233). It remains to be determined whether this subtle difference in positional distribution has any impact on the nutritional or health effects of 12:0 from the two sources. In any case, the suitability of transgenic plants for producing other medium chain fatty acids (8:0 and 10:0) is also being explored (234).

Finally, opportunities exist to use transgenic plants to produce waxes that may be designed to function as moisture barriers in foods. Potential industrial uses of waxes as lubricants and cosmetics will likely provide the impetus for developments in this area (148,212). However, an improved understanding of structure–function relationships between molecular characteristics of waxes and component migration in foods will place food scientists in a position to also benefit from wax production by transgenic plants.

C. Modifying Chemical Functionality

Genetic manipulation of lipid biosynthesis has already expanded to potential applications beyond the scope of storage lipids (triacylglycerols). The overexpression of Δ9DES in transgenic tomato (Lycopersicum esculentum) fruits results in increases
These transformations also enhanced the levels of pleasant character impact flavoring compounds [which are mediated by a lipoxygenase/hydroperoxide lyase/isomerase/reductase enzyme system (211)] in freshly macerated fruit tissues. Since tomato fruit has virtually no storage lipid, this study represents an approach to the modification of functional lipids in a manner designed to enhance their capacity as a reservoir for flavoring precursors.

The strategic manipulation of an assembly of lipids in the tomato fruit requires an understanding of polar glycerolipid assembly, which was beyond the scope of this chapter. (But the initial steps involve some of the glycerolipid assembly reactions in and beyond Fig. 3, and the Kennedy pathway in Fig. 5; functional glycerolipid assembly is covered in some detail in Refs. 15, 106, and 108.) Another potential extension of transgenic modification of membrane lipids is to enhance the postharvest storage life of plants tissues, particularly edible organs. Chilling sensitivity in plants and their organs is correlative to the proportion of phosphatidylglycerol (PG) that exists as species containing two saturated fatty acids (236). When plastidic GPAT from a chilling-insensitive species (spinach) and very chilling-sensitive species (squash, Curcurbita moschata) was cloned into a moderately chilling-sensitive species (tobacco, Nicotiana tabacum), tobacco constructs with the spinach GPAT had improved chilling resistance, whereas tobacco constructs with the squash GPAT had greater chilling sensitivity than did the wild-type tobacco plants (237). Relative chilling sensitivity of the transgenic plants was correlated to the proportion of disaturated PG. Thus, the variable reaction selectivities of plastidic GPAT among plants may have a central role in conferring chilling sensitivity by influencing the fatty acyl composition of polar glycerolipids assembled in the plastid. It should be noted that an alternative explanation for the aforementioned observations is that more global changes in lipid metabolism may account for the changes in sensitivity of transgenic plants to cool climates (16,238).

A more mainstream chemical functionality problem that may be ameliorated by genetic manipulation of storage lipid assembly is that of oxidative stability of plant oils. The most simplistic approach here would be to enhance the levels of monounsaturated fatty acids (18:1$_{9}$) and, concomitantly, to diminish the levels of polyunsaturated fatty acids (18:2$_{9,12}$ and/or 18:3$_{9,12,15}$) (ignoring for the moment the issue of essential fatty acids). Applications in this area were initiated with chemical mutagenesis in sunflower lines (239,240). High-18:1$_{9}$ sunflower oils have potential for use as oil sprays, dairy product substitutes, salad oils, and frying oils. Indeed, high-18:1$_{9}$ sunflower oil possesses greater thermal stability as a frying oil that does conventional sunflower oil (241).

Reductions in polyunsaturated fatty acid levels have also been achieved by genetic manipulation. In soybean and canola transformants, 18:2$_{9,12}$ has been reduced from 55% to 2.5% and from 19% to 6.3%, respectively, with a corresponding increase in 18:1$_{9}$ content (218). In this application, use was made of antisense technology toward genes encoding for Δ12DES (218,222). This approach was also extended to using antisense technology toward Δ15DES to reduce 18:3$_{9,12,15}$ content from 6.9% to 1.4% in transgenic canola (218).

A more complex and challenging, but still feasible, approach to engineering-improved oxidative stability relies on a thorough understanding of the influence of specific molecular species of triacylglycerols on the progress of lipid oxidation. The positional distribution of fatty acids (both saturated and unsaturated) along the glyc-
erol backbone of triacylglycerols can have a profound influence on the oxidative stability of an oil (242–244). If a systematic understanding regarding the influence of triacylglycerol molecular species on oxidative stability is developed (which arguably is primarily in the domain of food science), then the appropriate targets for genetic manipulation are easily identifiable. Unfortunately, elucidation of how triacylglycerol molecular species influence oxidative stability will take years, if not decades; yet such understanding is necessary before cogent genetic strategies can be identified to address the problem of oxidative stability within this context. The obvious advantages of genetically engineering oxidative stability are improved quality and stability of lipids in food, and potentially, a diminished dependence on exogenous antioxidants.

D. Modifying Nutritional Functionality

Some approaches to improving nutritional quality of plant-derived lipids involve genetic interventions just described (Sec. VI.C). Developing plant oils enhanced in monounsaturated fatty acids (18:1/9) are among these strategies, and the general health-promoting view of monounsaturates (239) has led to interest in developing high-18:1/9 lines for sunflower, corn, safflower, canola, and peanut oils. In addition, genetic manipulation to enhance Δ12DES and suppress Δ15DES would enhance levels of 18:2/9,12, arguably the most essential dietary fatty acid (213), although 18:3/9,12,15 is also essential (245,246).

Some microalgae (lower plants), such as Navicula pelliculosa, Cyclotella cryptica, and Phaeodactylum tricornatum, are capable of accumulating lipids enriched (18–26%) in 20:5/5,8,11,14,17 (247,248). Thus, the synthetic potential exists in the plant kingdom to yield 20:5/5,8,11,14,17 and 22:6/4,7,10,13,16,19, the two fish oil fatty acids (referred to as n-3, or ω3) that are widely believed to promote human health (246). Starting with 18:2/9,12, all the enzymes representing the sequence of steps involved in elongation and Δ6,5,4 desaturation, whether via the n-6 or n-3 pathway (246), have been demonstrated in higher plants: see Secs. III.B.6 (98,99), IV.C (167,168), and V.E.1 (193), as well as in algae (247,248) and fungi (4). Whether these traits can be incorporated in a manner that will permit them to collectively function to yield 20:5/5,8,11,14,17 and 22:6/4,7,10,13,16,19 in oils of transgenic plants depends on the abilities of the constitutive and cloned desaturases and elongases to coordinatively channel the acyl chain, alternatively linked to PC and CoA, toward these fatty acids, and then recognize them in triacylglycerol assembly steps. At least two immediate and specific concerns can be identified in this regard. Could the Δ4DES [of coriander: see Sec. III.B.6 (98,99)] be cloned in a manner effective at yielding 22:6/4,7,10,13,16,19 since the Δ4DES is believed to be plastidic stromal (soluble) enzyme acting on acyl-ACP, and its participation in 22:6/4,7,10,13,16,19 biosynthesis would be the terminal step, likely requiring action on a PC-linked acyl chain? Second, would the elongases, known to exist in Brassica and Crambe spp., as well as in leek, jojoba, and meadowfoam (see Sec. IV.B), be active with acyl with acyl chains of three to five double bonds, since only those of zero to two double bonds are elongated in vivo in wild-type plants?

Finally, opportunities exist to manipulate plants to yield “structured glycerides,” or triacylglycerol species that have specific nutritional functions that promote health (29,34,231) (see Chapter 28). The impact of dietary lipids on human cardio-
vascular health is a function not only of the fatty acid composition but of the positional distribution of the fatty acids along the glycerol backbone. An enhanced understanding of structure–function relationships of triacylglycerols and cardiovascular health would provide suitable targets for manipulating the pattern of triacylglycerol assembly in transgenic plants for the purpose of promoting health. A related example would entail producing medium chain triacylglycerols, which have value as dietary supplements for patients requiring medical and nutritional interventions (29). Other types of structured glyceride conferring nutritional benefit include Salatrim and Caprenin, which are mixtures of short and long chain fatty acids and contribute fewer calories to the diet than do conventional food oils (249) (see Chapter 22). Presently, these reduced-calorie fats are prepared by postproduction, chemical processes.

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REFERENCES
32. I. Ikeda, T. Oka, K. Koba, M. Sugana, and M. S. F. Lie Ken Jie. 5c, 11c, 14c-Eicosatrienoic acid and 5c, 11c, 14c, 17c-eicosatetraenoic acid of *Biot a orientalis* seed oil affect lipid metabolism in the rat. *Lipids* 27:500 (1992).


179. S. Stymne and A. K. Stobart. Evidence for the reversibility of the acyl-CoA: lyso-
phosphatidylcholine acyltransferase in microsomal preparations from developing sa-
180. R. Bernerth and M. Frentzen. Utilization of erucyl-CoA by acyltransferases from
developing seeds of *Brassica napus* (L.) involved in triacylglycerols biosynthesis.
181. G. Vogel and J. Browse. Cholinephosphotransferase and diacylglycerol acyltrans-
ferase. Substrate specificities at a key branch point in seed lipid metabolism. *Plant
182. B. Lemieux, M. Miquel, C. R. Somerville, and J. Browse. Mutants of *Arabidopsis*
in microsomal preparations from the developing seeds of safflower (*Carthamus tinc-
torius*) and turnip rape (*Brassica campestris*) and their ability to assemble cocoa-
184. G. Griffiths and J. L. Harwood. The regulation of triacylglycerols biosynthesis in
185. K. Ichihara. The action of phosphatidate phosphatase of the fatty-acid composition
186. G. Griffiths, S. Stymne, and A. K. Stobart. The utilisation of fatty-acid substrates in
triacylglycerol biosynthesis by tissue-slices of developing safflower (*Carthamus tinc-
188. M. Bafor, L. Jonsson, A. K. Stobart, and S. Stymne. Regulation of triacylglycerol
biosynthesis in embryos and microsomal preparations from the developing seeds of
189. M. Bafor and S. Stymne. Substrate specificities of glycerol acylating enzymes from
190. H. M. Davies, D. J. Hawkins, and J. S. Nelsen. Lysophosphatidic acid acyltransferase
from immature coconut endosperm having medium chain length substrate specific.
191. D. S. Knutzon, K. D. Laradizabal, J. S. Nelsen, J. L. Bleibaum, H. M. Davies, and
J. G. Metz. Cloning of a coconut endosperm cDNA encoding a 1-acyl-sn-glycerol-
3-phosphate acyltransferase that accepts medium-chain-length substrates. *Plant Phy-
192. J. F. Battey and J. B. Ohlrogge. A comparison of the metabolic fate of fatty acids of
193. M. R. Pollard and P. K. Stumpf. Biosynthesis of C<sub>20</sub> and C<sub>22</sub> fatty acids by developing
seeds of *Limnanthes alba*: Chain elongation and Δ<sub>9</sub> desaturation. *Plant Physiol.* 66:
649 (1980).
194. E. Fehling and K. D. Mukherjee. Biosynthesis of triacylglycerols containing very
long chain mono-unsaturated fatty acids in seeds of *Lunaria annua*. *Phytochemistry*
195. E. Fehling, D. J. Murphy, and K. D. Mukherjee. Biosynthesis of triacylglycerols
containing very long chain monounsaturated acyl moieties in developing seeds. *Plant
196. I. Löhden and M. Frentzen. Triacylglycerol biosynthesis in developing seeds of *Tro-


I. INTRODUCTION

A new oilseed crop was introduced commercially in the southern United States in the fall of 1994. The crop looked no different from normal varieties of Brassica napus canola. The farmer cultivated and harvested the crop without departing from standard canola harvesting practice. The oilseed meal, after crushing, was essentially the same as regular canola meal and, indeed, was treated simply as standard canola meal for use in animal feeds. The resulting vegetable oil, however, was unique and different from any previously available for either food or industrial uses (1).

The new oilseed crop was derived from transgenic canola developed by a biotechnology company. The genetic engineering approach was targeted toward a lauric acid rich oil in this case, but the technology is being used to develop other new vegetable oils with novel structure and compositions. In this chapter, we discuss some of the oils that may result and also some of the factors that shape the feasibility of these and other potentially novel raw materials to be achieved by genetic engineering.

II. GENETIC ENGINEERING OF PLANTS

The chemical composition of vegetable oils is a highly heritable trait. For example, year to year, soybean oil is a reliably constant raw material for the food industry. Not only is the high lauric acid content of coconut oil characteristic of coconut oil, but one does not find an occasional crop year in which soybean oil has lauric acid, or in which coconut oil is lacking lauric acid. These basic chemical compositions
characteristic of various vegetable oils—which define the uses of specific vegetable oils in the food industry—are determined by the genes of each plant variety. The advent of genetic engineering technology in agriculture has enabled the directed modification of the gene set that determines oil composition in a given oilseed crop.

There is a long and fruitful history of plant breeders who have deliberately selected for lines in which the seed oil is different in chemical composition. Examples include high-oleic sunflower, low-linolenic flax, and low-erucic rapeseed. These successes appear to be cases in which a specific gene or genes become nonfunctional. Although genetic engineering techniques in yeasts and bacteria allow the specific targeting of genes to be “knocked out,” this is not yet the case with plants. However, it is possible now to add genes to a plant, and some of these approaches can indeed be used to decrease the functional expression of genes resident in the host crop plant genome.

There is an ever increasing knowledge base to explain the genetic bases that determine the chemical composition of seed oils. In summary (2), a number of lipids-related genes have been individually cloned in the laboratory, and most of these genes turn out to encode specific enzymes used by the plant to synthesize the triacylglycerols that make up vegetable oils. For example, cloning a gene from Cuphea lanceolata that encodes a specific enzyme in the seed with a unique activity on a capric acid precursor may help to explain why high levels of capric acid are found in C. lanceolata seed oil. Moreover, failure to find that enzyme in canola seed may at least partly explain why capric acid is not naturally found in canola oil. And in its simplest manifestation, a genetic engineering approach would suggest taking the cloned C. lanceolata gene, transferring it into the chromosomes of a canola plant, and seeing whether the oil from the seeds of the resulting transgenic canola plant contains capric acid.

Currently, then, genetic engineering of plants allows the addition of genes. These genes are incorporated directly into plant chromosomes and basically behave in subsequent generations of progeny plant like other genes. That is, they are inherited in a Mendelian manner and are subject to the same modifications to which the genes preexisting in the plant genome are subject. The gene’s specific “behavior,” however, may be something entirely novel to the host plant. For example, it may encode for an enzyme that has never been found in that plant species. The ability to introduce such modifications is what genetic engineering adds to the plant breeder’s tool kit. A breeder, of course, could not cross a coconut tree with a canola plant to get a canola plant with certain coconut tree properties. But a discrete number of coconut genes can be transferred into a canola plant to make some facet of a canola plant’s metabolism more closely resemble that of a coconut tree.

As in the example of Cuphea lanceolata and canola above, one can add a novel gene from a wide range of sources, including not only any plant but also animals, bacteria, and even genes encoding enzymes that were “designed” on a computer and synthesized in the laboratory. It is possible to clone gene from a specific plant species, engineer the gene so that it expresses the same enzyme at an unusually high level at the targeted stage of seed development, and add it back to the same plant species. By having more of an enzyme controlling a rate-limiting step in oil biosynthesis, one might achieve more oil and/or an oil with a different fatty acid composition. And finally, it is possible to add genes that interfere with the function of
existing genes in the host plant genome, thus decreasing enzyme activities and enzymatic products that are not desired.

All these facets of adding genes can be used to redirect the molecular pathway by which oils are synthesized in plants; since these changes are determined by the genes, the changed oil in a transgenic plant will also be seen in the progeny of that plant. Thus, new crop species can be created that essentially look the same as the parent crop, yet produce a significantly different oil. A transgenic soybean plant looks like any other soybean plant and can be grown by the farmer in exactly the same ways.

A review of the detailed methods used to genetically engineer oilseed crops is outside the scope of this chapter. The reader is referred to several reviews (1,3,4). The intent of this summary is to describe some of the practical impacts on the food lipids area that will result from the introduction of a set of technologies that lead to new crop varieties, which in turn produce new raw materials for the food industry.

III. THE EXAMPLE OF LAURATE CANOLA

Coconut and oil palm kernel are the primary sources of lauric oils. Both are produced from trees grown in the tropical zone, and each contains about 50% lauric acid by weight. However, some temperate zone plants (e.g., Cuphea glutinosa, Umbellularia californica) accumulate even higher levels of lauric acid in seed oils. Therefore, a priori, it appeared in the early 1980s that lauric acid oils could conceivably be produced in temperate zone agricultural systems. For industries in the United States, a temperate zone crop source of lauric acid would (a) help address national import/export imbalances; (b) perhaps stabilize world price fluctuations for lauric acid oils; and (c) conceivably provide an oil with higher lauric acid content and the associated savings in processing costs.

When Calgene, Inc., of Davis, California, initiated a project in 1985 to engineer Brassica napus canola into a lauric oil producer, there was a long list of technical unknowns. There was no reliable system for putting genes into canola and getting back normal plants. The available data suggested that a gene from a monocotyledonous plant like coconut might not function correctly when transferred to a dicotyledonous plant like canola. The general opinion was that production of lauric acid in nonseed tissues might be detrimental or even lethal to a canola plant. Indeed, even as late as 1990, it was thought that lauric acid might “gum up” triacylglycerol synthesis in canola. Certainly in 1985 there was an insufficient understanding of how to limit foreign gene expression to just the developing seed of a transgenic canola plant. Also there was absolutely no experience, on which to base assumptions that a transgenic oil or simply the process of generating transgenic canola would not somehow compromise the agronomic productivity of canola. The perhaps most worrisome aspect of all was that no one knew, despite numerous tries, the mechanistic basis of laurate accumulation in coconut or oil palm kernel, let alone the wild species of Cuphea glutinosa or Umbellularia californica.

This list of technical challenges provides a framework to understand the steps and areas of technical expertise required to generate a useful new crop type containing a genetically engineered oil.

In the example of lauric acid canola, scientists were able to demonstrate the existence of a unique enzyme comprising lauroyl-acyl carrier protein (ACP) thio-
esterase in embryos of seed from *Umbellularia californica* that could conceivably account for the production of lauric acid. Moreover, that specific enzyme activity appeared to be missing in similar extracts from canola seed. Encouraged by that finding, a team of biochemists purified the *U. californica* lauroyl–ACP thioesterase protein sufficiently to obtain, by means of an automated protein sequencer, a partial amino acid sequence. This amino acid sequence allowed the design of synthetic DNA primers, which were then used with templates made by molecular biologists from messenger RNA isolated from developing seed of *U. californica* to generate gene-specific DNA probes. The DNA probes were subsequently used to identify lauroyl–ACP thioesterase complementary DNA (cDNA) clones from a cDNA bank that was constructed using, again, messenger RNA (mRNA) from developing seed of *U. californica*.

Independently in the same laboratory, another group of molecular biologists isolated a canola gene specifically expressed at high levels in developing embryos during the normal period of canola seed development when storage lipids are formed. This natural *B. napus* gene was dissected down to the elements necessary to encode proper gene expression timing and tissue-specific localization within the canola plant. These “promoter” or genetic expression elements were then combined with the central portion of the *U. californica* lauroyl–ACP thioesterase cDNA close corresponding to the open reading frame encoded by the original naturally occurring messenger RNA. This synthetic gene was then combined with a second gene (the selectable marker gene) in a specific manner relative to other DNA signal sequences in a unique microbe known as a disarmed *Agrobacterium tumefaciens*.

Plant cell biologists then took the genetically engineered strain of *A. tumefaciens* and cultivated it for a brief time with sectioned hypocotyl tissues from germinated *B. napus* plantlets. This now-routine method of gene transfer into canola had to be developed while the biochemists were studying the lauroyl–ACP thioesterase enzyme in *U. californica* seed extracts and while molecular biologists were identifying the gene expression control elements from embryo-specifically expressed genes in seed of *B. napus*. After the treatment of *B. napus* hypocotyl sections with *A. tumefaciens*, the bacteria were completely removed and the plant tissues cultured on a series of different growth media in different containers, to regenerate complete *B. napus* canola plants. During this process of regeneration, most of the plantlets were deliberately killed by addition of an antibiotic. The selectable marker gene that was linked to the synthetic lauroyl–ACP thioesterase gene in the *A. tumefaciens* strain was a synthetic gene designed to detoxify the antibiotic used in these experiments. Thus the only plants that survived the antibiotic treatment were transgenic plantlets that had received the kanamycin resistance gene (and almost always the lauroyl–ACP thioesterase gene as well). This is how the cell biologist can selectively produce 30 to 300 different transgenic plants without having to sort through thousands of plants that only might be transgenic. Each transgenic plant coming out of this process is potentially different and is generally regarded as a different event. As it turns out, each event tends to be unique in one or several features, most notably, in this case, with respect to how much lauric acid accumulates in the seed oil.

Progeny from each different event can be grown up and examined for the lauric acid content in the seed oil, and for other traits as well. As mentioned above, the transgenes show inheritance like regular canola genes, so gene segregation is possible, especially when a transgenic canola is crossed with another line.
It becomes the plant breeder’s job to genetically fix the transgenes in a given canola line as well as to select for other traits such that the lauric acid content is reliably constant from generation to generation and agronomic characters such as seed yield are also preserved. Practically speaking, of course, this means multiple seasons of field testing and selections. In addition, just like other canolas, lines optimally adapted to different environments have to be tested and selected in different environments. The importance of this phase to the eventual success of developing a new crop type with a novel oil composition should not be underestimated.

In summary, then, development of laurate canola required basic biochemistry research, applied protein purification, gene cloning molecular biology, embryo-specific gene expression molecular biology, cell biology, and breeding, with many of these tasks and much of the technology development carried out in parallel. As well as an agronomically acceptable lauric acid canola, there had to be development of an infrastructure at the farm level to produce and harvest a new canola without contamination by regular canola seed. And as we shall see, another set of expertises to recognize and take advantage of special properties that may be present in a new oil also had to be established. Clearly, the development of a new crop type by directed genetic modification of a seed oil is not a small undertaking.

IV. APPLICATIONS FOR FOOD LIPIDS

A. Natural Limitations

One of the first considerations in thinking of specific ways to modify seed storage lipids is feasibility. Certainly there are some limits to what is practically possible. For example, seed storage lipids serve an important role in the life cycle of a crop plant. Germinating seeds presumably require the energy stored in the seed lipids and/or the young seedlings require access to those seed lipids in the cotyledons. If a lipid has been modified into particular structures that interfere with the ability of germinating seeds or young seedlings to tap that energy, or if the oil contains fatty acids that are difficult to metabolize, there may be a limit to the quantities of modified lipids of those kinds that one can achieve in a seed oil and still have a viable crop variety.

Moreover, the synthesis and incorporation of certain unusual fatty acids into storage lipids during seed development may also result in those fatty acids becoming incorporated into structural lipids that are essential for normal seed function. For example, the castor bean endosperm contains very high levels of ricinoleic fatty acid in the storage triacylglycerols, with very little ricinoleic acid in structural lipids. Clearly, as the castor bean lineage evolved synthetic mechanisms for ricinoleic fatty acid in seed oils, it also evolved mechanisms to either prevent incorporation of ricinoleic acid into structural lipids or to clear that fatty acid from structural lipid molecules. If one chooses to engineer soybean to produce ricinoleic acid in seeds, it may be found that soybean lacks the ability to maintain the integrity of the seed structural lipids and the seed is not fertile.

It should perhaps be noted that lipid biosynthesis occurs in all plant cells and is essential to support growth. Whereas the triacylglycerols comprising vegetable oils found in seeds are neutral lipids used as a means to store energy and fixed carbon, polar lipids found in all cells—including those in seeds—serve important structural
functions and are thus often referred to as structural lipids. Both structural lipids and seed triacylglycerols contain glycerol-bound fatty acids, and thus it is not surprising that their respective synthesis pathways share many common steps. Re-engineering triacylglycerol biosynthesis must not compromise the synthesis of the structural lipids necessary to support growth and viability.

If some desirable lipid compositions are indeed incompatible with viable seed, there may be a unique opportunity in engineering such oil types into either oil palm or avocado mesocarp tissues. These are oil-rich tissues whose natural fate is to rot away after fruit dehiscence. Since plant progeny do not directly depend on the viability of these tissues, they might be engineered to “self-destruct” by making, for example, very solid fats rich in long chain saturated fatty acids. Engineered oil palm fruits could be harvested by standard means and the palm kernels (the oil palm seed) would be normal—assuming that the technology employed worked to ensure that the changed fatty acid composition was limited to just the mesocarp.

Another type of limitation that is of commercial interest is the ultimate amount of oil obtainable from a crop. While focused on traits like disease resistance and plant stature, much of the oilseed crop breeding community measures success by how much seed is harvested per acre. As crop yields go up, one expects the cost of food oils to come down. At a different level, breeders and genetic engineers alike are interested in increasing the oil content on a per-seed basis. Soybeans typically contain only 20–25% oil by weight, whereas canola seed is typically 40–45%. Peanuts can be 50% oil, while the cacao bean is 60% cocoa butter. It seems physiologically possible, therefore, to increase the seed oil content in soybean and canola with correspondingly dramatic effects on the cost of producing vegetable oils from these major crops.

B. Practical Limitations

Even though the scope of possible lipids modification projects must be very widely based on the natural variation in seed lipids found among plants form around the world, there are factors, other than natural ones, that practically limit what will be done in the near term.

Genetic engineering of crop plants requires a battery of expertises: plant physiology, enzymology, molecular biology, gene transfer cell biology, prototype evaluation, and plant breeding. Much of an engineering project entails sequential applications of each area of expertise to the eventual goal. Coupled to the obvious costs of having these expertises in place and having the time needed to go from concept to practice is a consideration of the technical risk entailed when one is embarking on a new project. Financial modeling and analyses of the eventual value and return versus development costs and risk may argue against ever starting certain projects.

Part of the eventual value may hinge on the production system that allows the value to be ensured and protected. Since genetically engineered genes behave like other genes once present in a plant, they are necessarily transmitted by pollen. Vegetable oils are often zygotic characters. That is, the chemical composition is determined by both parents. Thus if a transgenic canola crop is grown immediately adjacent to normal canola, an interchange of pollen will decrease the purity of both crops. So, novel oilseed crops may have to be grown in carefully managed districts. Of course, there are some precedents for producing “specialty” oils, including the
previously mentioned low-linolenic flax, high-oleic sunflower, and erucic-containing industrial rapeseed. These identity-preserved production practices typically add costs. Moreover, seed from crops with different types of oil cannot be stored together or crushed together; this restriction adds more complexity to production. These special production practices need to be considered when the development of a new food oil is being planned. In the case of canola, where the precedent of identity-preserved production of high erucic oils already exists, it appears that the incremental costs are likely less than $0.05/lb.

Regulatory and political issues should be anticipated as well. Since genetic engineering is a new technology, products are receiving additional scrutiny by the public and by regulatory agencies to ensure that environmental and food safety issues are as fully reviewed as feasible. The nature of the issues varies considerably according to the nature of the altered traits. For example, different issues are raised by herbicide resistance and the potential introduction of a new food oil rich in myristic acid. Some segments of society may be less willing to try foods containing ingredients based on biotechnology. Religious issues for some novel oils may need addressing (e.g., kosher and halal definitions). These market realities may affect what genetic engineering projects make sense.

Although plant lipid biosynthesis research has blossomed dramatically in the past 10 years, fueled by interest in transgenic plant applications, there is still much to be learned. Even though plant oils exist in nature that are greater than 85% laurate, there may not be enough known to justify the laying out of a scientific strategy to convert soybean into a producer of an oil with 85% lauric acid. On the other hand, if a technical route to some financially attractive target appears to be straightforward, an organization should consider the competitive aspects carefully. Perhaps a competitor elected to gamble at an earlier stage when the technical route was less clear and that organization may have gained an insurmountable lead to a commanding patent position.

Obviously, embarking on a long-term project with technical risk and complexity requires not only an evaluation of the utility and marketplace need of the final product, but also an extensive competitive analysis. The technology underlying the genetic engineering of oil composition may provide a clear basis for the patentability of the product of the research. Thus, since whoever is first may very well be in a position to block similar approaches by others, a consideration of which groups are after the same goal and where they are in the process is in order. In addition, for vegetable oils, there may be more than one means to the same end, and these alternatives should be anticipated as well as possible. For example, one could consider the genetic engineering of soybean with a *C. lanceolata* gene to produce an oil rich in capric acid. Alternatively, another group may choose more conventional means of domesticating *C. lanceolata* into an economically feasible crop. Either or neither approach may succeed; but if both succeed, the eventual marketplace value to each for the financial and time investment may be less.

In the case of structured oils and fats, there are of course chemically and enzymatically based synthetic methods that utilize glycerol and fatty acids sourced from animal fats as well as vegetable oils. Cost of raw materials and synthetic capacities are factors to consider when this approach is compared to the genetic engineering of crop plants. In the latter case, volumes are limited only to the number
of acres that can be planted; in the former case, capital-intensive hard assets may be required in the form of manufacturing plants.

Of course, the possibility of competitive products becoming available and concerns about the profitability and financial return on investment versus risk are critical concepts for the private sector entities considering the development of new food lipids via the genetic engineering of crop plants. On the other hand, certain targets of lipids modification that will not pass these private sector hurdles for profitability versus risk may nonetheless have value and importance to the human community. For example, a cottonseed oil lacking malvalic acid (see Sec. IV.B.4) may not be able to command much of a premium over regular cottonseed oil as long as cheap soybean oil (which naturally lacks malvalic acid) is also available. This might be a lipids modification project that could appropriately by undertaken by public sector research groups like the U.S. Department of Agriculture, since the benefit could be spread among a large number of independent cooperative ginning entities throughout the cotton-growing delta of the Mississippi River. Similarly, improving the nutritive value of lipids in rice may not be attractive to private sector companies for many reasons, including the long and perhaps risky research phases that would be necessary, the structure (or lack of structure) in how rice planting seed is sold, the prevalence of small (subsistence) family farms in specific growing regions of Asia, and the resistance of farmers to paying premiums. But clearly, if the technology can be used to improve traits like carotenoid or vitamin content even slightly in a basic commodity foodstuff, the benefits would be tremendous to society at large.

The important issue to address regarding public sector initiative research on tailored vegetable oils is the coordinated recognition of technical feasibility along with marketplace-defined needs (e.g., shelf life or processing traits like colors or free fatty acid content) and consumer common interest objectives (e.g., levels of atherogenic saturated fatty acids or essential dietary fatty acids). How this process sorts out common-good objectives and provides for the necessary funding and coordination of properly qualified laboratories, plant breeding organizations, and the equally necessary product development and introduction functions is a challenge for everyone working in areas relating to foods and health.

C. Target Crops

As of this writing, efficient systems exist to transfer genes into select tissue types of rapeseed (including canola), soybean, cotton, and corn; it is then possible to regenerate transgenic plants with normal growth habits and harvest yields similar to parental types. Thus, transgenic crops are already in commercial production (or there are existing seed sales) for transgenic canola, cotton, corn, and soybean.

Success has been reported by a few groups for similar gene transfer and regeneration steps for sunflower, flax, and peanut. The procedures for these and perhaps other temperate zone annual oilseed crops like safflower and sesame will likely become routine relative soon.

Because of the long generation times and the related lack of regeneration cell biology knowledge, current prospects for transgenic oil-producing tree crops like oil palm, coconut, olive, and cacao are less encouraging. Also of food interest is the modification of the fatty acids in oils of almond, walnut, and other nuts that tend to oxidize and produce off-flavors after storage. The time required with tree species to
develop and apply methods, evaluate transgenically produced potential products, and scale-up for production may discourage any commercial projects. Like the projects described above that do not offer large profit margin opportunities to reward the risk takers, genetic engineering of the lipids in major oil crops like oil palm and coconut may be taken up primarily by public sector research units, which do not need to satisfy short-term investment return expectations.

Another consideration in the modification of seed oil content of specific crops may be the ability to properly express transgenes in specific tissues of the plant. That is, seed lipid biosynthesis tends to occur during a very specific phase of seed development. To be able to modify that biosynthetic pathway to alter the character of the vegetable oil, transgenes must exert their effects in the correct tissue at the correct time. Moreover, in some cases, it may be deleterious to have a transgene expressed in the wrong tissue or at the wrong time. Ricinoleic acid naturally appears only in the castor bean endosperm and not in leaves and other tissues, where it presumably has no function and might interfere with the synthesis of necessary structural molecules.

Technology to obtain the correct transgene expression for seed storage lipid modification has been well demonstrated in canola and soybean. One can expect the same principles to apply for peanut, sunflower, and other crops, although fine-tuning of the technology may be needed for maximum benefit in some cases. This potential requirement should be taken into account in considerations of transgenic approaches to oil modification in crops where there is less experience.

**D. Target Traits**

1. **Short Chain Saturates**

The first genetically engineered vegetable oil, Calgene’s Laurical, is already a commercial product. Researchers studying the seed embryos of *Umbellularia californica* (California bay tree) identified an enzyme known as a lauroyl–ACP thioesterase that appeared to account for the 60% laurate content in the California bay seed oil (Fig. 1). When the cDNA corresponding to the mRNA for the bay tree lauroyl–ACP thioesterase was successfully cloned, adapted to canola gene expression controlling elements, and transferred into canola, the resulting oil from transgenic seeds contained lauric acid in amounts ranging from less than 1% to more than 45%. From these original plants, referred to as transgenic “events,” lines were developed that produced genetically uniform seed that reliably contained an average 38–42% lauric acid in the oil.

When the lauric acid rich oil from transgenic canola seed was examined more closely, it was observed that very little of the lauric acid was in the second position of the triglyceride molecules. The practical implications of this result are discussed below. From a scientific point of view, it was assumed that the canola enzyme that converts lysophosphatidic acid to phosphatidic acid discriminated against lauroyl-CoA as a substrate (Fig. 2). Indeed, it had been reported that in canola, this enzyme, known as lysophosphatidic acid acyltransferase (LPAT) discriminates in vitro against saturated acyl CoAs as well as substrates with monounsaturated acyl groups exceeding 18 carbons in length.

Subsequently, it was shown that coconut endosperm, which contains a triacylglycerol oil with high levels of laurate in the second position, also contains an LPAT
Figure 1  Biosynthesis of fatty acids in plant cells. (A) Synthesis of fatty acids in canola pro-plastids: fatty acids, or acyl groups, are built up in two-carbon increments while bound as a thioester to acyl carrier protein (ACP). As the acyl-ACP molecules lengthen into C16:0-ACP, C18:0-ACP, and (after a desaturation step) C18:1-ACP, an acyl–ACP thioesterase cleaves the thioester bond to ACP and the complete fatty acid is available for triacylglycerol synthesis after conversion to an acyl-CoA in the cytoplasm (see Fig. 2). (B) Engineered laurate biosynthesis in transgenic canola: addition of a lauroyl–ACP thioesterase from *Umbellularia californica* (or other lauric acid producing plant) allows the thioester bond to ACP to be cleaved at the C12:0 stage, thus making free lauric acid available for triacylglycerol synthesis.

Figure 2  Biosynthesis of triacylglycerols in plants. Reactions 1, 2, and 4 each draw on acyl-CoA pools available in the plant cell. (1) Glycerol-3-phosphate is acylated to form 1-acyl-sn-glycerol-3-phosphate, also known as lysophosphaticid acid. (2) 1-Acyl-sn-glycerol-3-phosphate is acylated to form phosphatidic acid by the enzyme lysophosphaticid acid acyltransferase (LPAT). (3) A phosphatase activity removes the phosphate group to generate diacylglycerol. (4) Diacylglycerol acyltransferase attaches the third acyl group to create triacylglycerol. LPAT in most plants is very selective for unsaturated 18-carbon acyl-CoA, whereas the other enzymes in this pathway are typically nonselective. However, coconut LPAT in particular has high activity on shorter chain saturated fatty acid CoAs.
enzyme with high activity for placing laurate into the second position. When the cDNA corresponding to the mRNA for the coconut endosperm LPAT was cloned, adapted to canola gene expression controlling elements, and transferred into canola along with the bay tree thioesterase gene construct, the resulting oil from transgenic seeds contained laurate in all three positions of the triacylglycerols. Thus, one type of lauric oil was obtained by engineering canola with the bay tree thioesterase enzyme, and a second type of oil was obtained by combining the bay tree lauroyl–ACP thioesterase with the coconut endosperm LPAT in seed of transgenic canola.

It has also been possible to create novel oils in seeds of transgenic plants with increased levels of other saturated fatty acids of chain lengths less than 16 carbons. By using cDNA clones derived from mRNA from different species of New World genus *Cuphea* plants, seed oils enriched in C8 and C10 fatty acids have been obtained. Using cDNA clones derived from mRNA from developing seed of *Cuphea palustris* or from nutmeg, oils enriched in C14 fatty acids have been obtained. The latter myristate-type oils also tend to have higher (C16:0 (palmitic) levels. Again, the transgenic canola triacylglycerols tend to have these newly introduced, engineered fatty acids in the first and third positions on the glycerol backbone. Thus, these oils and fats may have special functional properties due to the underlying structure.

2. Naturally Solid Fats

The temperate zone oilseed crops tend to have storage lipids rich in unsaturated fatty acids, and thus liquid oils that are in themselves not suitable for products like shortenings and margarine. This shortcoming in typically addressed by a postharvest treatment of partial hydrogenation of vegetable oils. Partial hydrogenation does carry some incremental cost and generally results in the introduction of trans-unsaturated fatty acids, which may be undesirable in foods for health reasons. Of the tropical tree oils, only oil palm is produced on a scale large enough to provide a “fat” fraction rich in palmitic acid as a hardstock. Illipe, shea, and sal are tropical trees with seeds rich in the C18 fatty acid stearate, which contributes more solids to a fat; however, such tropical fats are limited in availability and generally too expensive to use as hardstock ingredients. Inspection of the pathway for biosynthesis of fatty acids reveals that both palmitic and stearic fatty acids are precursors to the unsaturated fatty acids making up the bulk of the oil found in seeds from temperate zone crops. Not surprisingly, then, it has been possible to engineer the pathway so that fewer unsaturated fatty acids are made and more of the precursor palmitic and stearic acids accumulate in the triacylglycerols.

Numerous thioesterase enzymes have been identified by cloning cDNAs from plant mRNAs that share DNA homology with the California bay tree thioesterase cDNA described above. Every plant so far examined appears to have an enzyme with high activity on oleoyl–ACP as a substrate, and this enzyme presumably is the basic thioesterase in the fatty acid biosynthesis pathway shown in Figure 1. The oleoyl–ACP thioesterase typically has less but significant activity on both palmitoyl–ACP and stearoyl–ACP substrates. Another class of thioesterase enzymes has been found in many plants, however, with enhanced activity on palmitoyl–ACP substrates; genetically engineered overexpression of such enzymes in seed of transgenic canola plants results in seed oils enriched in palmitic acid. On the other hand, thioesterase enzymes with low activity on palmitoyl–ACP substrates and relatively high activity
on steroyl–ACP appear to be rare. When researchers looked at cDNA clones made from mRNA from developing seed of mangosteen fruit (mangosteen seed oil typically contains 40–50% stearic acid), a clone was identified that corresponds to an enzyme with enhanced levels of steroyl–ACP activity relative to other thioesterases. Genetically engineered overexpression of that mangosteen enzyme in seed of transgenic canola plants results in a seed oil enriched in stearic acid, up to 30% in some seed.

Alternatively, one can consider engineering a seed to have less desaturase activity so that fewer molecules of the saturated precursor steroyl–ACP are converted to oleoyl–ACP. This has been demonstrated in transgenic canola and soybean by suppressing levels of the steroyl–ACP desaturase enzyme. Suppression of an enzyme can be achieved by genetic engineering methods of either antisense or cosuppression. Each allows for tissue-specific suppression. In this particular application, tissue-specific suppression is important because the steroyl–ACP desaturase is an essential enzyme in leaves and other tissues for plant viability.

3. Yield

As noted above, yield is a very important trait in reducing the cost of vegetable oils. Theoretically, there are several arguments for the feasibility of raising seed oil content in most crops. The cacao bean has 60% oil by weight, so the physiological limit for crops like soybean and canola may be at least this high. Individual seed of rapeseed can typically vary from 35% to 50%; so even within existing germ plasm, the current average oil content from canola of 42% will likely be raised by straightforward breeding selections in the coming years. Historically, soybean protein content has been the most important component of the bean, with the 20% of the seed that is oil a valuable by-product. Clearly, there is room to increase oil content in soybean by some means; however, increased oil will be balanced against maintaining value in the meal component for animal feed uses.

Despite the evident technical premise and motivation for increasing oil content, exact scientific strategies are still developing. Acetyl–CoA carboxylase is an enzyme activity often considered to be a rate-limiting step for fatty acid biosynthesis. The data for this assumption seem clearest in animal cells and fairly convincing in the bacterium *E. coli*, but somewhat less clear in higher plants. Transgenic modifications of levels of this enzyme activity have been achieved with slight and possibly significant increases in oil content; however, experiments continue. The development of other strategies to increase fatty acid biosynthesis at the expense of nondigestible fiber and other less desirable components of seed represent an area of increasing research.

4. Removing Negatives

The availability of gene suppression technologies allows one to think of decreasing certain constituents of vegetable oils. An early and far-reaching application is evident in the use of cosuppression in canola and soybean to dramatically reduce the levels of polyunsaturated fatty acids. Other potential project might be, for example, the suppression of enzymes directly responsible for formation of cyclopropene fatty acids such as malvalic acid in cottonseed oil or achieving a decrease in the high palmitic acid content in palm oil by suppressing the level of palmitoyl–ACP thioesterase
activity in the oil palm mesocarp so that more fatty acids can be elongated and desaturated to oleic acid.

5. Other Lipid Targets

For the most part we have discussed modification of the kinds of fatty acid found in triacylglycerols. However, vegetable oils can also contain other important lipid components. These include the antioxidants of tocopherols (e.g., vitamin E) and of mixed carotenoids (including β-carotene and lutein), plant sterols such as sitosterol (which may have cholesterol-lowering properties in the human diet), and economically important by-products of vegetable oil processing such as lecithin. As the enzymatic bases for the synthesis of these compounds and the genetic bases for regulation of amounts of these compounds become better understood, there should be growing prospects for the genetic engineering of levels of these compounds as well as flavor components in vegetable oils.

E. The Nature of Incremental Progress

When low erucic acid lines of rapeseed were first commercialized in Canada, the oil content levels were typically less than 38% by weight. However, over years of breeding progress, oil content steadily increased to a current average of 42–44%. The primary determinant of value, oil quality, was first selected for and developed; then further incremental improvements in yield, meal quality, and other traits were layered onto the original trait.

Genetically engineered plant oils well each likely follow a similar path. That is, after the synthesis and accumulation of lauric acid have been established in a transgenic canola oil, it may become important to select (or genetically engineer) for lower levels of linoleic and linolenic fatty acids. Or as discussed above, a coconut LPAT enzyme may be engineered into the lauric canola to enable higher levels of laurate to be achieved. Such higher levels of lauric acid made possible by the presence of the medium chain LPAT enzyme may actually be achieved by field selections (breeding) over a number of years of incremental improvements.

And finally, of course, once a valuable oil has been achieved in composition, one would like to keep selecting for varieties with better and better yields. Thus, it is clear that genetic engineering of oil composition, just like conventional breeding of oil composition, comprises one or a few big steps—transgenes in the case of genetic engineering—supplemented by a continuous process of fine-tuning the oil and the crop variety producing the oil.

V. IDENTIFICATION OF UTILITY AND VALUE

A. Market-Driven Product Development

1. Fatty Acid Strategies

Fats and oils are ubiquitous components of most compounded formulations, whether they are foods or industrial products. These applications may not involve the whole triglyceride, or the pure derivative fatty acids that comprise triglycerides, but may entail the splitting of the triglyceride into its component fatty acids (see Fig. 3) and subsequent derivatization of those fatty acids into industrially important products that move into the oleochemical markets.
We shall not discuss these industrial applications at great length, but it is important to recognize their importance in the dynamics of the total fats and oils industry. Figure 4 is a typical process flow diagram showing the types of derivatives that are of current importance in the various industrial sectors. As can be seen from the end products synthesized and their uses, most fatty acids are converted to fatty alcohols and then to a variety of derivatives that utilize their surface activity. Although these compounds are, as a class, called surfactants, they perform a variety of functions in a variety of end products, including wetting agents, spreading agents, emulsifiers, foaming agents, and foam stabilizers. The majority of these surfactants are based on alcohols derived from lauric acid. Natural alcohols (from naturally occurring fats and oils containing C12:0 and/or fatty acids of interest >C12) compete with synthetic alcohols derived from petroleum. Synthetic alcohols historically dominated the market, primarily because of their low cost, but naturally derived alcohols are becoming increasingly important. Natural alcohol production is projected to equal that of the synthetics by the year 2000, as indicated by the data for fatty acid production in Table 1. No new synthetic alcohol capacity has come on-line in the last decade, and no new plants are projected for the near future. In contrast, six new natural alcohol facilities have been built since 1990.
After methylation of free fatty acids, the methyl esters are converted to fatty alcohols, which find their way into numerous consumer products. The driver for this industry is lauric acid availability, since this product yields derivatives having the most functional end effects. Natural lauric acid is available on a commercial basis only from the splitting and fractionation of certain tropical oils: coconut oil and palm kernel oil.

The world’s major coconut oil producer is the Philippines, where thousands of small farmers with small production acreage accounted for 43% of production and 65% of total exports in 1992–1993. Because the Philippine coconut industry has suffered from chronic underinvestment, production and exports have decreased and are expected to continue to do so for the foreseeable future. Estimates are that as

<table>
<thead>
<tr>
<th></th>
<th></th>
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<tr>
<td>United States</td>
<td>1300</td>
<td>1499</td>
<td>1653</td>
<td>592</td>
</tr>
<tr>
<td>Western Europe</td>
<td>1973</td>
<td>2226</td>
<td>2424</td>
<td>879</td>
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<td>Asia</td>
<td>1223</td>
<td>1455</td>
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<tr>
<td>Other</td>
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</tr>
<tr>
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<td>4915</td>
<td>5675</td>
<td>6303</td>
<td>2242</td>
</tr>
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</table>

Table 1 Volume and Value of World Fatty Acid Production
many as one-third of the 300 million coconut trees in the Philippines will become nonproductive over the next 10 years. Significant coconut oil producers are shown in Table 2. Production in these secondary countries, spurred by government investment, it is expected to increase. Coconut production in any given year is heavily influenced by weather factors, including rainfall amounts and the impact of typhoons.

The second major source of lauric oils is palm kernel. Malaysia is the largest producer of palm kernel oil accounting for 54% of the world’s production in 1992–1993. Other large producers are shown in Table 2. Unlike coconut production in the Philippines, palm (along with palm kernel) is produced by large sophisticated plantations and enjoys major government and private investments that are allowing it to forward-integrate into end-use derivatives. As a result, palm kernel oil production is projected to increase significantly through the year 2002, with yields of 2175 metric tons as new plantations become productive.

In the world of foods, where, typically the whole triglyceride is utilized as one of the components, lauric oils can be replaced—to some degree—by the nonlauric oils, especially as specialty fat blends are developed that mimic the physical properties of these lauric fats through a series of processing steps including special hydrogenation and fractionation. In the industrial area, however, lauric acid derived molecules invest unique properties into the surface activity of the resulting compound and formulation, and they allow for little or no substitution of the fatty acid moiety without a direct effect on performance. This is especially true in the soap, detergent, and personal care markets, where the C12 moiety provides unequaled detergency

Table 2  World Lauric Oils Production (million metric tons); By Country, 1992–1993

<table>
<thead>
<tr>
<th>Country</th>
<th>Coconut oil</th>
<th>Percent of world total</th>
<th>Palm kernel oil</th>
<th>Percent of world total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Production</td>
<td></td>
<td>Production</td>
<td></td>
</tr>
<tr>
<td>Cameroon</td>
<td>0</td>
<td>0</td>
<td>22</td>
<td>1</td>
</tr>
<tr>
<td>Colombia</td>
<td>0</td>
<td>0</td>
<td>31</td>
<td>2</td>
</tr>
<tr>
<td>Ecuador</td>
<td>0</td>
<td>0</td>
<td>11</td>
<td>1</td>
</tr>
<tr>
<td>Ghana</td>
<td>0</td>
<td>0</td>
<td>13</td>
<td>1</td>
</tr>
<tr>
<td>India</td>
<td>267</td>
<td>9</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Indonesia</td>
<td>664</td>
<td>23</td>
<td>336</td>
<td>20</td>
</tr>
<tr>
<td>Ivory Coast</td>
<td>47</td>
<td>2</td>
<td>31</td>
<td>2</td>
</tr>
<tr>
<td>Malaysia</td>
<td>32</td>
<td>1</td>
<td>925</td>
<td>54</td>
</tr>
<tr>
<td>Mexico</td>
<td>106</td>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Mozambique</td>
<td>39</td>
<td>1</td>
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<td>0</td>
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<tr>
<td>Nigeria</td>
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<td>0</td>
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<td>10</td>
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<tr>
<td>Papua New Guinea</td>
<td>37</td>
<td>1</td>
<td>21</td>
<td>1</td>
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<td>0</td>
</tr>
<tr>
<td>Sri Lanka</td>
<td>36</td>
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<td>0</td>
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<tr>
<td>Thailand</td>
<td>40</td>
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<td>27</td>
<td>2</td>
</tr>
<tr>
<td>Vietnam</td>
<td>107</td>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Zaire</td>
<td>0</td>
<td>0</td>
<td>22</td>
<td>1</td>
</tr>
<tr>
<td>Others</td>
<td>212</td>
<td>7</td>
<td>89</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>2888</td>
<td>100%</td>
<td>1704</td>
<td>100%</td>
</tr>
</tbody>
</table>
and mildness. Because the perennial nature of lauric oilseeds prevents quick response to changes in supply and demand, and much of the demand side is inelastic owing to the unique functionality of the oil, lauric oil prices are typically much more volatile than prices for nonlauric oils (Fig. 5). Rising incomes and standards of living in Southeast Asia and Eastern Europe are expected to result in an increase in consumption of detergent and personal care products and, therefore, increased demand for lauric oils. Nonetheless, most experts project that increases in palm kernel oil production will compensate for increases in laurate demand and decreases in coconut oil production. Lauric oil price volatility is expected to decrease from historic patterns, but prices are expected to remain much more volatile than those for the nonlauric products.

Under the circumstances just outlined, what better goal than to produce a temperate source of C12 fatty acids in a triglyceride using the techniques of genetic engineering coupled with traditional plant breeding? If high enough levels of C12 could be generated, several issues could be addressed:

1. The impact of wide fluctuations in commodity prices for world lauric oils due to weather, natural disaster, political unrest, and other uncontrollable factors could be mitigated by the development of an annual, temperate oilseed crop that could be used to offset the unpredictability of some of these tropical crops and to stabilize the pricing of the desired fatty acids.

2. Environmental concerns associated with the use of petrochemical products favor conversion to plant-oil-based products that are biodegradable and renewable. Chemical processing plants devote a significant portion of their capital to postproduction cleanup systems to minimize the environment stress. Use of plant oil feedstocks would ease environmental concerns and, presumably, reduce manufacturing costs.

The real world, however, imposes the following constraints:

![Figure 5](image-url)  
**Figure 5** Coconut oil prices: average price by year, in U.S. dollars per pound as crude oil delivered.
1. The basic oleochemical business is a coproduct-balanced, commodity operation. Profitability is dependent on tight control of manufacturing processes and costs, and on maximizing the value of every product produced as a result of the splitting process.

2. These industries are process driven. The plants have been designed around and optimized for profitability based on a relatively narrow range of feedstock composition. In the case of lauric fats, coconut, and palm kernel, the balance of the fatty acid streams produced dictates the profitability of the total operation. Major shifts in fatty acid distribution in the feedstock disrupt this balance and require changes in the way the end products are treated as saleable items. To clarify this point, note the difference in fatty acid composition of laurate canola versus palm kernel oil and coconut oil as shown in Table 3.

As can be seen, laurate canola does not contain as much C12 as either coconut oil or palm kernel oil. In addition to decreased levels of this fatty acid, note also the total absence of the medium chain fatty acids, C8:0 and C10:0, typically present in the tropical fats. Specific markets for these medium chains have been developed, and their sales contribution is important to the overall profitability of the splitting operation. The approach, then, to achieve utility within the existing framework of the massive oleochemical complex, would be to develop triglycerides having such massive doses of C12:0 that coproduct values would be obviated in standard feedstock sources. This would mean the development of triglycerides having C12:0 levels greater than 85%. Is such a goal possible utilizing this technology? The answers are not yet known, but the research still required is not trivial—as discussed above—and the impact of achieving such high levels of a specific fatty acid on the viability of the resulting plant is not known.

What has been learned from this exercise?

1. High expressions of specific fatty acids in plant triglycerides should be considered only in the context of alternative sources of such fatty acids, and the coproduct value of the rest of the components.

2. Supply limitations as drivers for specific fatty acids can be a viable foundation for product development if current sources of feedstock triglycerides do not necessarily offer cost offsets because of the value of coproducts.

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**Table 3**  Fatty Acid Composition (%) of Laurate Canola versus Coconut, Palm Kernel, and Canola Oils

<table>
<thead>
<tr>
<th>Fatty acids</th>
<th>Laurate canola</th>
<th>Coconut oil</th>
<th>Palm kernel oil</th>
<th>Canola</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lauric</td>
<td>38.0</td>
<td>49.0</td>
<td>47.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Myristic</td>
<td>4.0</td>
<td>17.5</td>
<td>16.0</td>
<td>0.1</td>
</tr>
<tr>
<td>Oleic</td>
<td>31.0</td>
<td>5.0</td>
<td>16.5</td>
<td>61.5</td>
</tr>
<tr>
<td>Linoleic</td>
<td>11.0</td>
<td>1.8</td>
<td>2.5</td>
<td>20.0</td>
</tr>
<tr>
<td>Other</td>
<td>16.0</td>
<td>26.7</td>
<td>18.0</td>
<td>18.4</td>
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<tr>
<td>Total</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
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</tr>
</tbody>
</table>

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In the context of developing increased levels of specific fatty acids in plant triglycerides, it is often fruitful to take a closer look at the actual coproducts derived from the feedstock splitting streams and to assess the value of those coproducts if they were not supply-limited (i.e., dependent on the levels present in the base feedstock oil). Again, coproduct value in the resultant genetically engineered oil must also be considered, so that value can be obtained using the whole product, and not depending on one or two of the fatty acids to carry the whole value of the product. Examples of potential products would include oils having very high levels of the medium chain fatty acids, C8:0 and C10:0, or very high levels of myristic acid, C14:0. Again, although these particular fatty acids are limited based on their presence in tropical oil feedstock streams, an assessment needs to be made of their potential value in the marketplace if such level-rich sources become available through genetic engineering. It is not a given that increased availability of these fatty acids would lead to an automatic increase in their industrial or food uses. One must also be constantly aware of the need to balance coproduct volumes and values to achieve success in the marketplace.

2. Triglyceride Strategies

An example of a strategy developed for the application of transgenic oilseed products is the use of laurate canola as an alternative for coconut and palm kernel fats (the so-called tropical laurates) in food systems. Although these markets are smaller in volume than the industrial applications of laurics (see Fig. 6), the functional nature of the uses would allow the recapture of the research investment, assuming that at least equal functionality was shown in existing applications.

The main function of lauric oils is to provide desirable structure and mouthfeel characteristics in a variety of foods. Both coconut and palm kernel provide relatively high solids at room temperature; but a steep melting curve at about body temperature gives food products based on these oils a good melt-away sensation in the mouth. The solids profile of a given fat is, typically, determined by dilatometric methods,

![Figure 6](image_url)

**Figure 6** World consumption of lauric oils by end use in 1994. Although food uses account for much of the consumption of lauric oils, oil where other oils with similar functionality could be used for cooking in most cases. That is, the low cost of lauric oils in developing countries is a primary factor in their use in food. However, soap and detergent uses, take advantage of the unique functionality associated with the 12-carbon lauric acid.
although techniques using pulsed NMR spectrometry allow for similar determinations of the solid/liquid ratio in a given fat as a function of temperature. Usually, the solids fat index (SFI) is determined for food fats at a select number of temperatures. The plots of solids versus temperature have been related over the years to the actual performance of given fat systems in specific food formulas. The portion of the curve below 25°C (77°F) relates to the “hardness” of the fat, specifically the resistance to deformation it provides to the base food, while the solids present between the temperatures of 25°C and 30°C (86°F) represent the amount of solids typically found at ambient temperatures and are a measure of the degree of heat resistance provided to the food product by the fat system. Solids that occur above 37°C (or about body temperature, 98.6°F) are perceived as “waxiness” in the mouth, although low levels of solids are acceptable. A rapid rate of solids decrease between ambient temperatures and body temperature provides a desirable cooling sensation in the mouth and is deemed a positive for confectionery fat systems. A major use of these fats is in confectionery coatings, where the lauric fats provide an effect similar to that of cocoa butter in chocolate, at a much reduced cost.

The major fat used in the manufacture of chocolate is cocoa butter. It is also one of the most expensive ingredients used in the formula, and because so much of it must be used to achieve the desired functional effects, it contributes the highest cost of goods for the total formulation. Aside from a significant contribution to the final flavor of the chocolate, cocoa butter has some physical properties that make it unique among the triglycerides found in nature. First of all, it is composed of a very few sets of highly structured triglycerides (see Table 4).

These triglycerides are essentially composed of only three fatty acids: palmitic (C16:0, P), stearic (C18:0, S), and oleic (C18:1, O). Practically all the oleic acid

Table 4  Triglyceride Composition of Cocoa Butter (%)

<table>
<thead>
<tr>
<th>Triglycerides</th>
<th>Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>POSt</td>
<td>36.3–41.2</td>
</tr>
<tr>
<td>StOSt</td>
<td>23.7–28.8</td>
</tr>
<tr>
<td>POP</td>
<td>13.8–18.4</td>
</tr>
<tr>
<td>StOO</td>
<td>2.7–6.0</td>
</tr>
<tr>
<td>StLiP</td>
<td>2.4–6.0</td>
</tr>
<tr>
<td>PLiS</td>
<td>2.4–4.3</td>
</tr>
<tr>
<td>POO</td>
<td>1.9–5.5</td>
</tr>
<tr>
<td>StOA</td>
<td>1.6–2.9</td>
</tr>
<tr>
<td>PLiP</td>
<td>1.5–2.5</td>
</tr>
<tr>
<td>StLiSt</td>
<td>1.2–2.1</td>
</tr>
<tr>
<td>OOA</td>
<td>0.8–1.8</td>
</tr>
<tr>
<td>PPSst</td>
<td>8.0</td>
</tr>
<tr>
<td>PSiSt</td>
<td>0.2–1.5</td>
</tr>
<tr>
<td>POLi</td>
<td>0.2–1.1</td>
</tr>
<tr>
<td>OOO</td>
<td>0.2–0.9</td>
</tr>
</tbody>
</table>

A, arachidate; Li, linoleate; O, oleate; P, palmitate; St, stearate.
occurs esterified at the sn-2 position of the various triglycerides, with the sn-1 and sn-3 positions containing the esterified palmitic or stearic acids. This specificity results in the presence of three predominant structured triglycerides: POP, POS, and SOS. These three triglyceride types make up more than 80% of the triglycerides found in cocoa butter. These specific triglycerides, which resemble each other very closely, provide cocoa butter with its unique solids profile and desirable melting characteristic.

Going back to the basic issue, however, cocoa butter is expensive, and supply limited. Over the years, various researchers have spent considerable amounts of time developing fats that could serve as alternatives to cocoa butter by providing properties similar to its melting profile and solids content at various temperatures. Several classes of alternatives have been developed, ranging from cocoa butter “equivalents” produced from selected blends or fractions of natural fats high in specific triglyceride contents that are miscible in all proportions to cocoa butter, to fats that contain totally different triglyceride distributions but mimic, in many ways, the melting behavior of cocoa butter. An example of the production of a cocoa butter “equivalent” would involve the purification and fractionation of a series of different naturally occurring fats to obtain the proper proportion of the desired triglycerides having the desired structure, at the appropriate levels. Commonly used sources of these specialty fats and their triglyceride distributions are given in Table 5. The majority of the POP portion required is obtained from palm midfraction.

The reconstruction of a cocoa butter equivalent from the isolation of specific triglycerides, through fractionation from a variety of source oils and subsequent blending, obviously is not an inexpensive process, especially since the oils used for feedstocks are sourced from tropical plants that are not grown and cultivated in the most efficient manner and in themselves are often supply-constrained. These “equivalents” are commercially viable only when the price of cocoa butter is high enough to justify their high costs. When cocoa butter is relatively inexpensive, they are not used.
Another approach to mimicking the functional properties of cocoa butter is through the use of special fractions of a combination of domestic and tropical (non-lauric) fats that have been selectively hydrogenated. The fats often used include soy, canola, cotton, and palm. The manufacture of chocolate-flavored coatings using these final blends of fat in place of cocoa butter is well known, and these coatings have a real place in the market in certain applications where rate of melting of the fat in the mouth and ultimate flavor release are not key factors in consumer acceptance of the product. They do have the drawback, however, of some amount of “waxiness” that can be detected in the mouth, since a fraction of their triglycerides does melt above body temperature. This effect is minimized when chocolate-flavored coatings made with these cocoa butter replacers are used on baked goods, where the crumb structure of the substrate aids in the mastication of the coating and helps minimize the perception of waxiness in the mouth.

The most popular approach for the formulation of chocolate-flavored coatings is the use of cocoa butter substitutes (CBSs), based on lauric fats. These coatings are widely used in both the chocolate/confectionery and baking industries. The principal base fats used in their manufacture are palm kernel oil and coconut oil, with the former being the more widely used. The specific fatty acid composition of these fats and their various fractions after processing is shown in Table 6. These fats contain triglycerides that are high in the esters of lauric and myristic acid. Triglycerides of these compositions form crystals that are relatively small and uniform, and have a sharp melting point around body temperature. The set point of these fats is also, typically, very sharp, and they provide a coating base that has significant solids at room temperature. With their high levels of lauric and myristic acids, however, it is obvious that a significant percentage of these fatty acids will occur on all three of the available positions, in a random manner. Because these triglycerides are of vastly different compositions and are nonstructured, coatings based on them are not compatible with cocoa butter; hence the ultimate coatings can tolerate only minimal levels of cocoa butter containing ingredients, such as cocoa powder, and the flavor impact of these coatings is usually degraded by these restrictions. They do have better melting properties than the cocoa butter replacers discussed above, however, with the noticeable absence of any lingering waxy mouthfeel.

With the production of the first crop of laurate canola, we were faced with a totally new type of triglyceride: one that was high in lauric acid esters, though not

<table>
<thead>
<tr>
<th>Oil</th>
<th>C12</th>
<th>C14</th>
<th>C16</th>
<th>C18</th>
<th>C18:1</th>
<th>C18:2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Palm kernel oil</td>
<td>47.0</td>
<td>17.0</td>
<td>8.5</td>
<td>3.0</td>
<td>11.0</td>
<td>2.0</td>
</tr>
<tr>
<td>Hydrogenated at 35°C</td>
<td>47.0</td>
<td>17.0</td>
<td>8.5</td>
<td>13.0</td>
<td>7.0</td>
<td>—</td>
</tr>
<tr>
<td>Hydrogenated at 40°C</td>
<td>47.0</td>
<td>17.0</td>
<td>8.5</td>
<td>19.0</td>
<td>1.0</td>
<td>—</td>
</tr>
<tr>
<td>Fractionated</td>
<td>55.0</td>
<td>21.0</td>
<td>8.5</td>
<td>2.0</td>
<td>7.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Hydro/fractionated</td>
<td>55.0</td>
<td>21.0</td>
<td>8.5</td>
<td>10.0</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Interesterified/hydrogenated</td>
<td>47.0</td>
<td>17.0</td>
<td>8.5</td>
<td>19.0</td>
<td>1.0</td>
<td>—</td>
</tr>
<tr>
<td>Coconut oil</td>
<td>47.5</td>
<td>17.5</td>
<td>8.5</td>
<td>2.5</td>
<td>7.0</td>
<td>1.5</td>
</tr>
<tr>
<td>Hydrogenated</td>
<td>47.5</td>
<td>17.5</td>
<td>8.5</td>
<td>10.0</td>
<td>1.0</td>
<td>—</td>
</tr>
</tbody>
</table>
as high as coconut or palm kernel oils, but one in which all the C12:0 and C14:0 occurred on the sn-1 and sn-3 positions, and the sn-2 position was occupied solely by C18:x fatty acids, where x = 1, 2, and/or 3. Selective hydrogenation of this oil, then, would allow us to closely control the melting properties of the resulting triglyceride and to manipulate the solids profile of the final fat system. When these new fats were utilized in typical formulated food systems, it was noted that the systems offered flavor release that was far superior to that obtainable with the corresponding tropical laurics. Although this statement is based on anecdotal data, a great deal of effort is under way to quantify the flavor release properties of these new structured fats versus the random lauric fats typically used. It could be immediately demonstrated, however, that this new source of a lauric fat used in confectionery was highly compatible with cocoa butter—a trait not encountered with the tropical laurics—hence could be used with cocoa butter sources that lead to higher flavor in the finished product. These obvious differences in how the genetically engineered laurate canola performed with respect to existing laurics were explored at some length and are discussed later (Sec. V.B.2).

3. Investigations Driven by Purported Health Benefits to the Consumer

a. Low Saturates. Consumer concern over saturates has driven food marketers to look for ways to differentiate their products on the basis of saturated fat content. Food labeling regulations recently published by the U.S. Food and Drug Administration (FDA) allow products containing less than 3.4% total saturates to be labeled as containing no saturated fat. With the size of the current salad and frying oil markets as a target, the potential justified examining and developing a technical strategy to address this opportunity. Over 45% of the vegetable oil consumed in the United States is used in salad and cooking oils. More than 6.5 billion pounds of vegetable oils (valued at $1.7 billion, wholesale) was used in this segment in 1992. The U.S. salad and cooking oil market is characterized by slow but steady growth, driven largely by population increases. The principal oils used in the production of retail salad and cooking oils in the United States are soybean, canola, corn, sunflower, cottonseed, and peanut oils (see Fig. 7).

Saturate levels in liquid oils have become a significant issue for consumers because of reports linking saturates to coronary heart disease. Saturates have been shown to lower the levels of high density lipoprotein (HDL) cholesterol. The FDA now requires that food labels list saturate levels. These concerns have already had a major effect of the U.S. salad and cooking oil segment, driving canola oil (with lower saturates than other oils) consumption from 263 million lb in 1987 to over 1.2 billion pounds in 1993. Consumption of saturated tropical fats decreased by 17.8% over the same period. Consumer concern over saturates is projected to continue to play a key role in this market segment, and food companies will continue to search for products that address this concern.

Since canola oil contains the lowest level of saturates of any of the commonly used food oils, it seemed a natural base from which to launch a variety of approaches that would lead, ultimately, to the desired product. As it turned out, the research was not trivial. A combination of mutagenesis and the use of two different genes led to a canola oil having a saturates level between 4.0% and 4.5%, but with questionable agronomic performance. The presence of relatively high levels of polysaturated C18 fatty acids also contributed negative performance characteristics when the oil
was used for frying purposes. Several different companies are continuing to work in this project area using conventional plant breeding to lower the saturates. In most cases, these companies are working to combine this attribute with low linolenic and/or high oleic traits to offer an oil that is low in saturates with improved stability under high temperature uses. None of these groups has yet developed an oil with less than 8% saturates.

**b. High Saturates.** The presence of *trans* fatty acids in the diet has become a major issue for U.S. and European consumers. In 1993 margarine sales began to decline as a number of studies indicated a potentially adverse impact of the *trans* fatty acids found in partially hydrogenated vegetable oils used in the manufacture of most margarines and spreads. These studies (some of which received extensive publicity and are discussed in Ref. 5) suggested that *trans* fatty acids contributed to coronary heart disease by raising levels of LDL cholesterol and lowering levels of HDL cholesterol. Although there is still considerable debate in the medical community about the true health impacts of *trans* fatty acids, they have become a source of concern for many consumers and have affected buying habits. At the same time that U.S. consumers became concerned over the *trans* issue, butter manufacturers began to aggressively compete with margarine manufacturers for market share through price reductions and increased marketing and promotional activities. The net result is that margarine sales have declined since 1993, falling by more than 6% from 1993 to 1994.

An opportunity was seen to exist for the development of a suitable oil containing sufficient saturates in its natural makeup to eliminate the need for hydrogenation. Hydrogenation, normally used to increase the saturated fatty acid levels in a triglyceride system, is the primary producer of *trans* fatty acids when polyunsaturated systems are reduced to monounsaturated systems. Thus elimination of hydrogenation would result in the elimination or minimizing of *trans* fatty acid occurrence in food.
systems. Key to assessing the opportunity is understanding the needs of the margarine and spreads industries.

Margarines and spreads are prepared by blending fats and oils with other ingredients, including water, milk, edible proteins, salt, flavorings, coloring. Spreads are differentiated from margarines by their lower fat content. By FDA regulation, a product must contain at least 80% fat to be labeled margarine; products with lower fat levels must be labeled as spreads. As consumers have become more concerned about overall fat consumption, spreads have become increasingly popular. The key to the formulation of both margarines and spreads is to provide a solid, spreadable fat with appropriate melting characteristics. This is achieved by using partially hydrogenated vegetable oils, which have had a portion of their unsaturated fatty acids converted to both saturated fatty acids and less saturated, trans fatty acids. Saturated and trans fatty acids have higher melting points than unsaturates and provide the requisite functional properties for the finished product. The specific oils used and the degree to which they are hydrogenated vary as a function of specific marketing objectives for the particular product.

In Europe, some margarines contain an oil portion consisting of a liquid vegetable oil blended with one that has been fully hydrogenated—a formulation ploy seen increasingly in the United States. Unlike partially hydrogenated oils, fully hydrogenated oils do not contain any trans fatty acids. This allows for the manufacture of margarines that are trans-free but also contain higher levels of saturates than those containing partially hydrogenated oils. European manufacturers also use more tropical oils in the manufacture of margarines and spreads than firms in the United States, since European consumers are not as concerned as U.S. consumers about tropical fats.

As of this writing, several companies are actively pursuing the development of seed oils that contain levels of saturated fatty acids high enough to permit the elimination of needs for hydrogenation, with the concomitant production of trans fatty acids. As an integral part of this higher production of functional, saturated fatty acids, it is also necessary to have the desired fatty acids in a low polyunsaturated fatty acid background, to assure the ultimate stability of the finished formulation. To compete in the large markets involved under these two sets of conditions of composition, one must also strive for a resulting plant with agronomic vigor and oil yields comparable to those of existing sources of commodity oils.

B. The Discovery of Novel Utility

1. Conventional Wisdom Versus Structured Triglycerides

Most of the body of knowledge that has been built up over the years on the functionality of lipids in food systems has been based on experiments designed around variations of naturally occurring fats and oils. The functional performance of these fats, from whatever source, was related back to specific analytical characteristics that still enjoy wide use in the industry: solid fat index, iodine value, and fatty acid composition. In addition to these characterizing values, a number of analytical tests were routinely performed on the fats that were indicators of their quality, or their ability to withstand the stresses of temperature and shelf-life requirements. These included free fatty acid content, peroxide value, color, and odor. None of these tests, however, related the functional performance of the fat to the presence (or absence)
of any specific triglyceride having a specific structure (i.e., which fatty acid was on which carbon of the glycerol backbone). In the majority of cases, such knowledge would have been of academic interest only, since these structured fats were simply not available in any great proportion in a given fat system. Aside from cocoa butter and some of the other more exotic tropical fats, most fats used in foods consist of a random assortment of triglycerides driven by the types and levels of fatty acids in their composition, so that such knowledge would have no direct bearing on a formulator’s capabilities.

Although many pioneering studies were conducted on synthesized and purified structured triglycerides to ascertain their physical chemical properties, especially those related to their melting characteristics and crystal forms, the quantities synthesized were not sufficient to be utilized in real food systems.

Recently, a great deal more effort has been expended to study structured triglycerides in foods, and these researches have led to the market introduction of synthesized species that have been almost exclusively targeted at the confectionery market for the replacement of cocoa butter, with the additional benefit of producing reduced-calorie products. These products take advantage of the effects of positional isomerism on the glycerin backbone to address the specific physical properties required in the final food product, and to utilize the differences in caloric contribution of the various fatty acids used to arrive at a lowered caloric intake. These novel ingredients, however, are costly to manufacture. Each one requires a series of synthetic steps along with requisite purification procedures. With a final price to the end user that remains at several dollars per pound, the ultimate use of novel ingredients is restricted to specific niche markets in the food industry; a significant move toward their use in a wide array of food products cannot yet be projected.

It is the foregoing type of research, however, that is needed to drive our understanding of the functionality of triglyceride structures in food systems. With the advent of the tools provided by genetic engineering, the opportunity to create new, structured triglycerides in the seed oil of an agricultural crop at costs much closer to a commodity seed oil base than to that of a synthesized product is very real. The goal, then, is to develop knowledge that relates structure to function so that a specific structure can be utilized for a specific end use. When this has been achieved, it is also likely that the total amount of fat required in any given food system will be reduced to levels well below those currently required using the various random systems as they occur in nature. The end result will very likely take us to the reduction in total fat intake that is so strongly recommended by health care professionals.

2. Laurate Canola: A Case Study

The first genetically engineered oil approved for food use was developed by Calgene over a period of approximately 10 years, through the use of techniques discussed earlier. The product was ultimately brought to market using the common and usual name, laurate canola. The specific composition of this oil was given earlier (Table 3). Initial functional screening of this new oil was conducted to see if it would serve as a cocoa butter replacement in coating and confectionery products. Although laurate canola did not contain the same levels of laurate as the coconut and/or palm kernel products currently used for these applications, all the C12 and C14 fatty acids occurred exclusively at the sn-1 and sn-3 positions of the glycerin molecule, and this
property encouraged us to look for novel functional effects. Since the \( sn\)-2 position contained only C18:\( x \) fatty acids, where \( x = 1, 2, \) or 3, selective hydrogenation would allow us to vary the SFI profiles of the resulting fats in a significant manner. Prior to commercializing a specific product, we were able to study a variety of plant oils that had increasing levels of the C12/C14 substitution, and to examine the effects of hydrogenation on the resultant SFI curves of the final fats. These data are shown in Figure 8.

The slope of the SFI curve clearly becomes steeper as the levels of C12 increase. It is also apparent that the tailing of the curve toward the higher melting end (the so-called waxy portion of the curve) is minimized as the C12 content increases. If we examine a specific laurate canola having approximately 38% C12 content, and vary the degree of hydrogenation, (i.e., vary the content of C18:0 vs. the C18-unsaturates that occupy the \( sn\)-2 position), we find the effects on the SFI curves illustrated in Figure 9.

As a reference, Figure 9 also provides the SFI for palm kernel stearine. The solids profile is bracketed by two laurate canola products: one with an IV of 17 and one with an IV of 37. Thus we knew that we could effectively match the solids profile, at least in the melting range around body temperature, that most manufacturers required for their products. The next most important piece of information for

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**Figure 8** Solid fat index (SFI) of laurate canola as a function of C12 content (wt %): squares, 30/32%; diamonds, 36%; circles, 40%. All values for laurate canola hydrogenated to an iodine value (IV) of 37.

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the use of this new fat in confectionery was knowledge of its crystallization properties. Most lauric fats used in this application crystallize in the $\beta$ form, without the need to go through any elaborate tempering step during the processing of the coating mass. Through X-ray crystallography, we were able to show that laurate canola crystallizes predominantly into a $\beta'$ crystal. With this knowledge, we then evaluated the product in a standard confectionery coating formulation (Table 7) versus com-

![Figure 9](image.png)

**Figure 9** Solid fat index (SFI) of cocoa butter, palm kernel stearine, and Laurical: 38% laurate canola as a function of IV (circles, IV 17; triangles, IV 37) compared to palm kernel stearine (diamonds) with an IV of 7 and cocoa butter (squares) with an IV of 34.

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sugar, 6×</td>
<td>49.6</td>
</tr>
<tr>
<td>Cocoa powder, natural (10–12)</td>
<td>11.0</td>
</tr>
<tr>
<td>Nonfat milk powder</td>
<td>8.0</td>
</tr>
<tr>
<td>Laurical 25</td>
<td>28.0</td>
</tr>
<tr>
<td>Laurical 15</td>
<td>3.0</td>
</tr>
<tr>
<td>Lecithin</td>
<td>0.3</td>
</tr>
<tr>
<td>Vanillin</td>
<td>0.1</td>
</tr>
</tbody>
</table>
commercial lauric fats based on both PKO and coconut oil. The results of these experimental evaluations revealed some significant differences between the laurate canola coatings and those made using palm kernel oil- or coconut oil-based fat systems, including:

- Significant increased flavor impact with laurate canola
- Increased coating shelf life (decreased bloom) when laurate canola was used
- Preferred mouthfeel over standard laurics

There were some negative findings also, including:

- Decreased shrinkage on cooling (problems with demolding)
- Less snap (coating not as hard as comparable controls)
- Less gloss (reflectance of surface crystals below that of controls, indicating that crystal size was an issue)

All these negative issues are being addressed through a combination of formulation techniques and procedures, and through variations in the structural makeup of the laurate canola itself. We believe that this fine-tuning process will be an integral part of any applications development effort for any of the structured triglycerides developed using this technology.

One of the most significant differences we found for the laurate canola versus the typical laurics used in confectionery coatings was the high degree of compatibility between laurate canola and cocoa butter. Typical lauric fats have limited compatibility with cocoa butter and tend to produce eutectic effects in admixtures that create softer fats than either of the two base fats alone. This result is primarily due to the interference of the crystallization path of the lauric fat on the crystallization dynamics of the cocoa butter. The addition of only a few percent of cocoa butter into a typical lauric fat will result in this softening effect, causing the resulting fat blend to be unsuitable for use in coating applications. With laurate canola, however, such negative interactions with cocoa butter did not occur until significant levels of cocoa butter (about 40% on an oil basis) were admixed. This means that sources of “chocolate” flavor, typically those high in cocoa butter, can now be used to impart more of the desired flavor to the finished goods when laurate canola is used as the base fat for the coating. The actual mechanism of this cocrystallization effect has not been determined, but the functional effects of such blends, as interpreted through SFI curves of the laurate canola–cocoa butter systems, is clear.

Again, finding advantages such as these and relating them to the composition and structure of the base oil and then using breeding and selection to “grow” the optimal oil will be ongoing efforts. However, once a baseline of information has been developed that relates structure to function, a predictive capability should be established that will significantly shorten the turnaround time required for new product development.

VI. PROSPECTS AND SUMMARY

We have seen that genetic engineering of dramatically new vegetable oil compositions is feasible, through a complex and protracted process leading to an economically viable transgenic crop that is then optimized by a continuing phase of improvements to that new crop variety. The fatty acid composition of vegetable oils is plastic;
moreover, new lipid structures that take advantage of those fatty acid building blocks are also possible.

The complexity and cost of practicing this technology and the concomitant planning imply a careful examination of the value and utility of the resulting vegetable oils before a genetic engineering project is begun. However, the potentially most exciting applications are those that create truly novel oils. Such oils typically do not have any extensive history of use in the food industry, and thus there are only small knowledge bases available to help predict value and utility of hypothetical oils. Certainly in the case of laurate canola, the significant performance improvements based on the unique triacylglycerol structure could not have been anticipated because that structure had not been predicted. Industry laboratories cannot justify spending resources to develop application for raw material vegetable oils they cannot obtain. Conversely, biotechnology companies will have trouble justifying the development of a novel oil if no industry group has demonstrated interest in buying that oil.

There is a long history of working with the established and conventional vegetable oils such as soybean or cottonseed, and therefore a certain familiarity with how they can be used. When truly novel vegetable oil and fatty acid compositions are made available, a careful reexamination of the basic assumptions for how commodity vegetable oils are currently used will be worth while. Important new applications may be possible due to inherently novel chemical composition and structures of vegetable oils created by the modification of crop plant genes. This, then, is a major challenge to lipid chemists in all fields: to understand the contributions of underlying composition and structure of oils to the eventual functionalities, and to use that understanding to intelligently predict the most advantageous uses of genetic engineering technology applied to vegetable oils.

REFERENCES