INCI Nomenclature

The goal of the INCI Nomenclature Program is to develop unique, informative, standardized and globally accepted names (called INCI Names) for the label declaration of ingredients used in cosmetic and personal care products. INCI Names are published in the *International Cosmetic Ingredient Dictionary and Handbook* which is the authoritative worldwide reference of ingredient information for industry, government, consumers, academia and the medical community. The International Nomenclature Committee (INC) is charged with the responsibility of creating the INCI nomenclature system, and assuring the integrity of the data incorporated into the *Dictionary*. The guiding principles below outline the approach followed by the INC in developing INCI Names. These conventions have evolved since the onset of ingredient labeling in the late 1970s, and continue to be revised as new ingredient innovation and technologies emerge. Central to the development of an INCI name is ingredient composition. Safety and suitability of the intended use of an ingredient is not reviewed as part of the INCI process.

INCI NOMENCLATURE CONVENTIONS

The conventions used to determine INCI names for cosmetic ingredients are listed below and are divided into three areas: *General Conventions*, *Specific Conventions* (which are grouped primarily by chemical class), and *Miscellaneous Conventions*. These conventions are continually reviewed and modified when necessary to reflect changes in the industry, technology, and new ingredient developments. Every effort is made to ensure ingredients are named consistent with these principles. As new conventions are developed that give rise to INCI names that are different from those previously published, the older nomenclature is sometimes retained and considered to be “grandfathered”. Grandfathered names are generally published for reference only.
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February 2021
GENERAL CONVENTIONS

1. Nomenclature assignments are based on the chemical composition of the intended product, and simple chemical names are used when possible. The assigned names are generally based on an ingredient’s final composition and purity irrespective of the type of manufacturing process (e.g., chemical synthesis, biotechnology, etc.). An ingredient is considered a single constituent or well-defined substance in accordance with the criteria outlined in the Technical Guidance Document for identification and naming of substances in REACh. Exceptions include organic and silicone polymers, botanical extracts, fermentation products and minerals, which are typically named by their starting materials and can include the process by which they are manufactured. For additional information, see the Conventions for Botanicals, Minerals, Polymers, and Silanes and Siloxanes.

2. Recognized chemical abbreviations are used where applicable. A list of the abbreviations used in the Dictionary may be found in the List of Acronyms.

3. Traditional stem names are retained as combining terms when consistent with other conventions. Commonly recognized trivial names will be utilized where appropriate, e.g., acylated derivatives of amino acids, Lauroyl Lysine, or acylated derivatives of hormones, N-Caffeoyl Serotonin. See also Rules 27a and 57.

4. Name/number combinations are used as INCI names for cosmetic ingredients only where the complexity and/or similarity of ingredients precludes assignment of reasonable nomenclature. The stem names are suggestive of the structure or the composition of the material, e.g., rh-Oligopeptide-6, Quaternium-27, Polyurethane-5, Polysilicone-1. Where descriptive terminology is desired for a particular component of a raw material that would fall under these classifications, alternate nomenclature may be provided. Established name/number or name/acronym combinations are also utilized, e.g., Red 4, CI 10020, HC Blue No. 10, Ceramide EOP.

5. Specific names previously established by the U.S. Pharmacopoeia (USP), National Formulary (NF), the Food Chemicals Codex (FCC), Merck Index, International Non-Proprietary Names for Pharmaceutical Substances (INN), World Health Organization (WHO), the Research Institute for Fragrance Materials (RIFM), and United States Adopted Names (USAN) are retained in many cases. Furthermore, established abbreviations and criteria are utilized for simplifying the nomenclature of families of complex ingredients where applicable. For example, the root word “alkonium” from the USAN convention to denote N,N-dimethyl N-alkyl benzyl in Benzalkonium Chloride is utilized to name other similar cosmetic ingredients such as Cocoalkonium Chloride. Compounds that are similar to materials described in recognized sources are given analogous names when possible. Examples include the Research Institute for Fragrance Materials (RIFM) names for the fragrance compounds, Linalool, Longifolene.

6. USAN abbreviations and criteria are utilized for simplifying the nomenclature of families of complex ingredients where applicable, e.g., Poloxamer, Nonoxynol, Octoxynol, Meroxapol.
7. Compounds that are similar to materials described in recognized sources are given analogous names when possible, e.g., Cocoalkonium Chloride.

8. Names of ingredients that contain terminal numbers are generally hyphenated, (see Convention 4), and names for derivatives of hyphenated materials retain the original hyphenated term, e.g., Quaternium-18 Hectorite.

9. Hydration states are not usually expressed, with the exception of Hydrated Silica. For information on process terms, see Convention 73.

10. Compounded mixtures created by blending materials are named by listing each component in descending order of predominance. See the section on Labeling Reminders for further information. On a case-by-case basis, materials containing a significant amount of unintended product may be named as a mixture, e.g., excess glycerin in products obtained by the transesterification.

11. Water, ethyl alcohol and other common diluents or solvents contained in commercially available raw materials, except extracts and products derived by fermentation, have not historically been identified as part of the INCI Name. In recent years, the INC has included the identification of non-aqueous solvents in an effort to provide further transparency. See the Labeling Reminders section for additional information on the labeling of solvents and/or diluents that may be present in raw materials.
   a. The INCI name for water, regardless of source is Water. The only exception is the INCI name Sea Water because it is sufficiently different in composition from fresh water, and it also corresponds to an EU Trivial Name, Maris Aqua. (See the Trivial Names section). Botanical waters are named in accordance with the Botanical Conventions.
   b. Purity standards or processes associated with water are not identified in the INCI name, e.g., purified water, deionized water, sterilized water.

12. For products marketed in the United States, the phrase “and other ingredients” may only be used in the label declaration when confidentiality has been granted by the U. S. Food and Drug Administration in accordance with procedures established in 21 CFR 701.3 and 720.8(a). Currently, there are no similar exclusions in the European Union Regulation (EC) No 1223/2009. For products marketed in Japan, the phrase “and other ingredients” may only be used in the label declaration when confidentiality has been granted by MHLW in accordance with procedures established in MHLW Notice No.990 (Sept/29/2000).

13. In order to facilitate clarity and ease of use when labeling, INCI names have been designed to require a minimum of punctuation and capitalization. slashes are used to designate compounds (not blends) that are produced as a mixture or are composed of more than one entity, e.g., Dipentaerythrityl Dicaprylate/Caprate, Glyceryl Cocoate/Citrate/Lactate, Styrene/MA Copolymer, Silicon/Cerium/Titanium/Zinc Oxides. Additionally, slashes are used to identify botanical materials that are hybrids. The terms are described in alphabetical order, separated by a slash. For additional information, see Convention 21 on the identification of alkyl groupings, and Convention 44(b) on the listing of monomers.
14. Wherever new nomenclature has been adopted, every effort has been made to use the shortest name consistent with these rules. Shorthand abbreviations will be considered for names with exceedingly long character length.

15. The International Cosmetic Ingredient Nomenclature Committee, in conjunction with the Personal Care Products Council, reserves the right to provide specific nomenclature in certain cases to make the nomenclature more informative to the consumer. In particular, terminology for ingredients related to drug active substances may be retained, (e.g., see Convention 50, Prostaglandin Derivatives).

SPECIFIC CONVENTIONS

Alkanolamides

16. Alkanolamides are named by the specific alkyl amide stem and the appropriate abbreviation, e.g., “Acetamide MEA,” “Lauramide DEA,” “Cocamide DIPA.”

Alkoxylated Materials

17. Alkoxylated materials are named by including the alkoxylation level as the average number of moles of ethylene oxide and/or propylene oxide, and/or ethyleneimine. Ethoxylates, propoxylates, and ethyleneimine are commonly expressed by the degree of polymerization. Mixed alkoxylated ethers (i.e., contain both EO and PO) are named based on the order of addition. For example, PPG-y Glycereth-x means glycerin is first treated with x moles EO then y moles PO. PEG/PPG-x/y means that a material is treated simultaneously with EO and PO, e.g., PEG/PPG-52/32 Dimethyl Ether. PPG-x-PEG-y indicates that the material is first ethoxylated, and then propoxylated.

18. a. Ethoxylated alcohols are named by adding the suffix “eth” to the conventional stem name followed by the average number of moles of ethylene oxide, e.g., Steareth-10. Historically, where the moles of ethoxylation is 1,2, or 3, the numerical designation is sometimes omitted, e.g., Sodium Laureth Sulfate, and the definition specifies the average number of moles as 1 to 3.

b. The term Alkoxynol-n refers to an ethoxylated alkyl phenol where n is the average ethoxylation value e.g., Nonoxynol-10. The following table references the alkoxynol stem to its alkyl group:

<table>
<thead>
<tr>
<th>Alkoxynol</th>
<th>Alkyl Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Octoxynol</td>
<td>Tetramethylbutyl</td>
</tr>
<tr>
<td>Nonoxynol</td>
<td>Nonyl</td>
</tr>
<tr>
<td>Dodoxynol</td>
<td>Dodecyl or Tributyl</td>
</tr>
<tr>
<td>Pentadoxynol</td>
<td>Pentadecyl</td>
</tr>
</tbody>
</table>

19. a. The polyethylene glycol fraction of all ethoxylated compounds not named as above is abbreviated as the acronym “PEG.” This combining form is followed by the average
number of moles of ethylene oxide. When the ethoxylating agent is 2-chloroethanol, names are generally designated by the term “hydroxyethyl.” See Convention 38.

b. Polypropylene glycol is abbreviated as the acronym “PG.” This combining form is followed by the average number of moles of propylene oxide, e.g., PPG-24 Butyl Ether. When the propoxylating agent is 2-chloropropanol, names are generally designated by the term “hydroxypropyl.” See Convention 38.

c. Polyethylene imine is abbreviated as the acronym “PEI”. This combining form is followed by the average number of moles of ethylene imine (aziridine), e.g., Hydroxyethyl PEI-10.

d. Homopolymers of ethylene glycol and propylene glycol are named as PEG-X and PPG-Y, respectively, with X or Y equal to the average total number of moles of alkoxyate in the material. Homopolymers of aziridine are named as PEI-X, with X equal to the average total number of moles of ethylene imine in the material.

e. Alkoxylated esters are named as PEG and PPG derivatives, e.g., PPG-10 Stearate, PEG-40 Stearate. Mixed alkoxylating esters are named in the order of addition of the alkoxylating agent. An example of the random simultaneous addition of the alkoxylating agent is e.g., PEG/PPG-8/3 Diisostearate, PEG/PPG-10/2 Ricinoleate. An example of the ordered sequential addition of the alkoxylating agent is PEG-4 PPG-13 C13-15 Alcohol.

f. PEG and PPG polymers or their derivatives in which one of the terminal primary alcoholic groups (CH2OH) has been oxidized to the carboxy group (-COOH), are named by adding the terms “carboxylic acid” or “carboxylate” to the parent name of the original polymer, e.g., Steareth-10 oxidized to carboxylic acid would be named Steareth-10 Carboxylic Acid.

g. Poloxamers, Meroxapols, Poloxamines and Minoxapols are named in accordance with convention 6 above. The term “Poloxamer” denotes a block copolymer consisting of polypropylene glycol terminated with polyethylene glycol. The term “Meroxapol” denotes a block copolymer consisting of polyethylene glycol terminated with polypropylene glycol. The term “Poloxamine” denotes a block copolymer of ethylene diamine reacted first with polypropylene glycol and then polyethylene glycol, e.g., [(PEG)_{x}-(PPG)_{y}]_{2}-NCH_{2}CH_{2}N-[(PPG)_{y}-(PEG)_{x}]_{2}. “Minoxapol” is the reverse of “Poloxamine”, e.g., [(PPG)_{x}-(PEG)_{y}]_{2}-NCH_{2}CH_{2}N-[(PEG)_{y}-(PPG)_{x}]_{2}. The numerical suffix designation is obtained by the following rule: The first two digits multiplied by 100 correspond to the approximate average molecular mass of the poly(oxypropylene) portion; the third digit multiplied by 10 corresponds to the percentage by weight of the poly(oxyethylene) portion.

h. Block and random copolymers of polyethylene glycol and polypropylene glycol not named in 19g. are named as PEG-X/PPG-Y Copolymer (block), and PEG/PPG-X/Y (random) where X is the average ethoxylation value and Y is the average propoxylation value, e.g., PEG/PPG-240/60 Copolymer. The sequence (block or random) and the terminal groups are described in the monograph definition of each ingredient.

i. Terpolymers with a center anchor, in which there is further block or random
alkoxylation of an alkoxylated polymer, are named, for example, as PEI-y PEG-x/PPG-y Copolymer. (e.g., PEI-14 PEG-10/PPG-7 Copolymer)

j. Ethoxylated glycerin is named as “Glycereth-x” where x denotes the average moles of ethylene oxide. Esters formed by the reaction of a fatty acid with an ethoxylated glycerin molecule are named by adding the suffix “ate” to the fatty acid grouping, e.g., Glycereth-5 Cocoate. When glycerin is derivatized prior to ethoxylation, (e.g., esterified with a fatty acid), the ethoxylation is designated by PEG-x, e.g., PEG-7 Glyceryl Cocoate, PEG-3 Glyceryl Trioleate.

Alkyl Groupings

20. The nomenclature for ingredients which are inherent mixtures (e.g., unfractionated fatty acids, or fatty alcohols from natural oils) is determined on the basis of the chemical identity of the raw material as purchased, (i.e., source and purity). Inherent mixtures that reflect the original distribution of components (i.e., when there is no fractionation) are named according to the common name of the source, e.g., Coconut Acid, Coconut Alkane, Soy Acid, Tallow Alcohol. Derivatives of these materials are named in a similar manner, e.g., Ammonium Palm Kernel Sulfate, PEG-5 Avocadoate, Tallowaminopropylamine. If the original natural distribution has been significantly cut or enriched, the mixture is named on the basis of the predominant component, e.g., Sodium Myreth-3 Sulfate. The predominant component was historically defined as one that is clearly present at the highest concentration in relation to the other components. Current practice defines the predominant component as a single constituent present at 80% or more of the composition. See Convention 1.

21. Nomenclature for materials that result from feedstocks that are mixtures, and where a single component does not predominate, (e.g., mixtures of fatty acids), is designated by the names of the major alkyl groups that make up 80% of the composition separated by a slash, e.g., Caprylic/Capric Glycerides, Glyceryl Isostearate/Myristate, Pentaerythrityl Stearate/Caprate/Caprylate/Adipate,. Coconut/Palm Kernel Alkanes, Coco/Sunfloweramidopropyl Betaine, Palm/Stear/Behenamidoethyl Diethonium Hydrolyzed Wheat Protein. When a mixture is constituted by a broad range of alkyl groups, “C” type nomenclature is used to designate a name, e.g., C14-30 Alkyl Beeswax, C10-16 Alkyl Glucoside. See Conventions 22 and 39.

An exception to this convention is the historical usage of the terms “cetearyl” and “cetoleyl” to identify a feedstock mixture of cetyl/stearyl alcohol and cetyl/oleyl alcohol, respectively. Historically, the name Stearic Acid has been applied to materials commercially sold as stearic acid where the composition is predominantly C16. Also, the term “vegetable” has been historically applied to a limited number of ingredient names, e.g., Vegetable Oil, Hydrogenated Vegetable Glycerides, Hydrolyzed Vegetable Protein, and their derivatives, and these names have been “grandfathered”. Current conventions stipulate the identification of the specific oil source(s) in the name, see Convention 43a.

22. Materials containing mixtures of even-carbon, straight-chain length fractions in which there is a predominant component are named by the common name for the predominant fatty stem. Materials containing mixtures of even- and odd-carbon chain length fractions are designated by alternative nomenclature when there is not a predominant component. Historically, the latter names were identified as “pareth”, e.g., C12-15 Pareth-3. The current
The nomenclature approach utilizes “alketh” as a stem name, e.g., C12-15 Alketh-3. See Conventions 21 and 39.

23. Straight-chain alkyl groups are described by their common stem names. The following table describes the nomenclature applied to straight-chain acids and alcohols.

<table>
<thead>
<tr>
<th>Chain Length</th>
<th>Acid</th>
<th>Alcohol</th>
</tr>
</thead>
<tbody>
<tr>
<td>C6</td>
<td>Caproic</td>
<td>Hexyl</td>
</tr>
<tr>
<td>C7</td>
<td>Heptanoic</td>
<td>Heptyl</td>
</tr>
<tr>
<td>C8</td>
<td>Capryl</td>
<td>Capryl</td>
</tr>
<tr>
<td>C9</td>
<td>Pelargonic</td>
<td>Nonyl</td>
</tr>
<tr>
<td>C10</td>
<td>Capric</td>
<td>Capryl</td>
</tr>
<tr>
<td>C11</td>
<td>Undecanoic</td>
<td>Undecyl</td>
</tr>
<tr>
<td>C12</td>
<td>Lauric</td>
<td>Lauryl</td>
</tr>
<tr>
<td>C13</td>
<td>Tridecanoic</td>
<td>Tridecyl</td>
</tr>
<tr>
<td>C14</td>
<td>Myristic</td>
<td>Myristy</td>
</tr>
<tr>
<td>C15</td>
<td>Pentadecanoic</td>
<td>Pentadecyl</td>
</tr>
<tr>
<td>C16</td>
<td>Palmitic</td>
<td>Cetyl</td>
</tr>
<tr>
<td>C17</td>
<td>Margaric</td>
<td>Heptadecyl</td>
</tr>
<tr>
<td>C18</td>
<td>Stearic</td>
<td>Stearyl</td>
</tr>
<tr>
<td>C20</td>
<td>Arachidic</td>
<td>Arachidyl</td>
</tr>
<tr>
<td>C22</td>
<td>Behenic</td>
<td>Behenyl</td>
</tr>
</tbody>
</table>

Unsaturated:

<table>
<thead>
<tr>
<th>Chain Length</th>
<th>Acid</th>
<th>Alcohol</th>
</tr>
</thead>
<tbody>
<tr>
<td>C11</td>
<td>Undecylenic</td>
<td>Undecylenyl</td>
</tr>
<tr>
<td>C16</td>
<td>Palmitoleic</td>
<td>Palmitoleyl</td>
</tr>
<tr>
<td>C18</td>
<td>Oleic</td>
<td>Oleyl</td>
</tr>
<tr>
<td>C18</td>
<td>Linoleic</td>
<td>Linoleyl</td>
</tr>
<tr>
<td>C18</td>
<td>Linolenic</td>
<td>Linolenyl</td>
</tr>
<tr>
<td>C20</td>
<td>Arachidonic</td>
<td>Arachidonyl</td>
</tr>
<tr>
<td>C22</td>
<td>Erucic</td>
<td>Erucyl</td>
</tr>
<tr>
<td>C22</td>
<td>Cetoleic</td>
<td>Cetoleyl</td>
</tr>
</tbody>
</table>

24. Branched-chain alkyl groups are described in INCI names by the prefix “iso” followed by the common stem name for the comparable straight-chain group. In such cases, usage of the term “iso” implies the isomeric nature of the carbon chain, (i.e., the same number of carbons in a nonlinear structure) rather than methyl substitution on the second carbon (i.e., in the iso position).

25. The following table has been included to clarify the nomenclature for derivatives of caproic, caprylic, and capric acids.

<table>
<thead>
<tr>
<th>Chainlength</th>
<th>Stem Name</th>
<th>Acid</th>
<th>Ester</th>
</tr>
</thead>
<tbody>
<tr>
<td>C6</td>
<td>Capro</td>
<td>Caproic</td>
<td>Caproate</td>
</tr>
<tr>
<td>C8</td>
<td>Capryl</td>
<td>Capryl</td>
<td>Caprylate</td>
</tr>
<tr>
<td>C10</td>
<td>Capr</td>
<td>Capric</td>
<td>Caprate</td>
</tr>
</tbody>
</table>
Amphoteric Compounds

26. The term “ampho” has been used as a combining term in the nomenclature for amphoteric surfactants derived from imidazoline intermediates. In naming these compounds, “ampho” denotes N-hydroxyethyl ethylenediamine and is combined with the names for the substituent groupings, e.g., Sodium Cocoamphoacetate.

Biological Materials (Excluding Botanicals and Biotech Materials)

27. Biological materials are named by a specific component (e.g., Hyaluronic Acid, Phosphatidyl Choline, Sphingosine) when the material has been isolated, purified and chemically characterized. General nomenclature for biological materials (e.g., Glycosaminoglycans, Fish Serum Extract) is utilized to name materials in accordance with the extent of their purification.

a. Trivial names for biological compounds are generally utilized in INCI names rather than systematic nomenclature, e.g., lysine, melatonin, lecithin. Trivial names are also used for derivatives where possible, e.g., N-Feruloyl Serotonin, N-Nicotinoyl Dopamine, Palmitoyl Arginine. See also Convention 3.

b. Ingredients derived from human tissue contain “human” as part of the INCI name, e.g., Human Umbilical Extract. See also Convention 61g.

c. Materials derived from animal sources are named, in most cases, by the common name of the animal, rather than the genus/species of the animal, e.g., Donkey Milk. Genus/species information may be included in the definition. For mammalian derived cells, the name of the organ is typically used unless a specific cell type has been isolated, e.g., Liver Cell Extract, Leukocyte Extract, Human Keratinocytes,

d. INCI names for fungi are identified by genus and species, e.g., Mucor Circinelloides Oil, Ganoderma Lucidum (Mushroom) Extract. Historically, the term “mushroom” is also included in the relevant INCI names since it is a term recognized by the consumer. Yeast has historical usage as an INCI name and is a grandfathered term. However, products derived by yeast fermentation are named by the genus term “Saccharomyces” (see Convention 28b.)

e. Yeast has historical usage as an INCI name and is a grandfathered term. It is retained in publication for its correlation to the EU Trivial Name, Faex. Products derived by yeast fermentation are named by the genus term “Saccharomyces” (see Convention 28b.)

f. Extracts of algae have historically been named as Algae Extract, in addition to being named by genus terms, e.g., Ascophyllum, Laminaria, etc. These names are grandfathered; the current naming practice is to include both the genus and species terms in the name. The INCI name “Plankton” is also a grandfathered term, and the
current practice is to also identify these materials by genus/species.

g. Coral, jellyfish, and sea anemone are invertebrates of the phylum, *Cnidaria*, and are named by their common name in accordance with Convention 28(c). Additional information about the classification of these species is included in the monograph definition when available.

h. Proteins are named by the common name, e.g., Collagen, Elastin, Keratin, Albumen, etc. Historically, protein materials that have undergone complete hydrolysis were named as amino acids by source, e.g., Keratin Amino Acids, Silk Amino Acids. These names have been grandfathered and the current naming practice does not distinguish by the extent of hydrolysis, e.g., Hydrolyzed Collagen, Hydrolyzed Milk Protein, Hydrolyzed Soy Protein.

i. Gelatin is one of the original names published in the Dictionary and is considered to be a “grandfathered” term. Usage of Gelatin as an ingredient name has been replaced by Hydrolyzed Collagen.

**Biotech Materials and Ferments**

28. Biotechnology involves the usage of microorganisms, or single cells in culture, for the industrial production of a material. Diverse ingredients may be produced through biotech processes, and distinct names may be assigned when the bioengineered material differs significantly from its traditional counterpart. (See also Synthetic Peptides, Convention 56h-k, for the nomenclature of peptides derived by recombinant technology.)

Biotech materials may be derived by the action of microorganisms, such as bacteria or yeasts, on a substrate to produce materials by fermentation, metabolism, hydrolysis, lysis, or other process. The process may involve the use of nutrients and other materials such as enzymes. The resulting product is referred to as a “culture” or “ferment.” The “ferment” may be further processed by extraction, filtration, or other procedure to yield the final product. The conventions used to provide INCI Names for biotech materials are as follows:

a. When the end product produced from a given “ferment” or “culture” has a common or usual name, such name may be used, e.g., Yogurt, Gellan Gum, Xanthan Gum, Wine.

b. When the end product does not have a common or usual name, the product is named using the genus of the microorganism followed by a slash, and the name of the substrate (if applicable), followed by the word “ferment.” Typical fermentation substrates (i.e., glucose, peptone) are not included as part of the INCI name. On a case-by-case basis, the genus and species name of the microorganism may be used when the use of the genus only may be misleading and the species is needed for clarity, particularly where pathogenic organisms are involved, e.g., Candida Bombicola Ferment, Escherichia Coli/Glucose Ferment Filtrate, Bifida/Enterococcus Faecium/Lactobacillus/Streptococcus Thermophilus/Soy milk Ferment Curd. Substrates will be identified by their common, usual, or other technical name, e.g., Lactococcus/Carrot Ferment. In the absence of a common name, the substrate may be named by the Latin genus/species term, e.g., Aspergillus/Camellia Sinensis Leaf Extract Ferment.
c. If a component(s) of the fermentation process has been isolated and purified to a significant extent (e.g., >80% based on dry weight), and analytical evidence is provided, the name for one or more of the components may be used, e.g., Glycosphingolipids, Beta-Glucan, Dextran, Sodium Hyaluronate.

d. Products derived by fermentation and further processed by extraction or filtration are named accordingly, e.g., Lactobacillus/Oat Ferment Extract Filtrate. When cells are lysed by mechanical treatment, heat treatment, osmotic pressure, by the use of chemicals, or enzymes, the term “lysate” is included in the name, e.g., Lactobacillus Ferment Lysate.

e. Conditioned Media is the growth media collected from eukaryotic cell cultures and are named according to the source of cells being cultured, e.g., Human Cord Blood Cell Conditioned Media, Camellia Sinensis Callus Culture Conditioned Media. No distinction is made regarding the growth conditions or manufacturing method, (e.g., cells grown under hypoxia or induced by ozone). Further distinction is given to indicate the presence of stem cells when the material is confirmed by appropriate analytical methodology. See 28f.

f. The naming of materials derived from stem cells is based on species and source tissue and is contingent upon full analytical characterization of the cell population using appropriate current methodologies, e.g., FACS, flow cytometry with appropriate markers and gating strategy. The INCI name is assigned based on whether the product is the media, the cells, or both. Examples include Horse Adipose Stromal Cell Conditioned Media, Human Bone Marrow Stem Cell Conditioned Media, Human Fibroblast Induced Pluripotent Cell Extract, Human Umbilical Mesenchymal Stem Cell Conditioned Media, Red Junglefowl Induced Pluripotent Cell Culture Conditioned Media, Sheep Adipose Stromal Cell Conditioned Media.

g. Exosomes are small vesicles found in the cells of eukaryotic organisms. With appropriate analytical characterization, (e.g., electron microscopy, flow cytometry, nanoparticle tracking analysis) they are named according to species and source tissue, e.g., Human Adipose Stromal Cell Exosomes, Human Amniotic Fluid Induced Pluripotent Cell Exosomes.

     Plants can make small, nano-sized, vesicles which are a structure found within or outside of a cell, consisting of liquid or cytoplasm enclosed by a lipid bilayer. These vesicles have not been characterized to the extent as those derived from mammalian cells, therefore they are named as vesicles. They are named based on the plant source that they are isolated from, e.g., Brassica Oleracea Italica (Broccoli) Vesicles, Dendropanax Morbiferus Leaf Vesicles. If these vesicles are released outside the cells then they are considered extracellular vesicles and are named based on the plant source they are released from, e.g., Panax Ginseng Adventitious Root Extracellular Vesicles, Rosa Damascena Callus Extracellular Vesicles.

h. Ingredients derived by plant tissue culture are named according to their method of production as follows:

     i. “Plant Callus Extract” is an extract of a callus, or any plant cells that are forced to go through a callus stage in culture unless re-differentiated, (see ix. below)
“Plant Callus Extract Powder” is an extract of callus cells which have been dried, and ground

“Plant Callus Culture Extract” is an extract of a whole culture (cell + media) of callus cells.

“Plant Callus Powder” is a callus (or callus cells), that has been dried and ground without extraction.

“Plant Cell Culture” is a cell suspension obtained using high shear and enzymatic treatment and without a callus formation.

“Plant Cell Culture Extract” is the extract of a whole cell suspension culture (cell + media).

“Plant Cell Culture Conditioned Media” is the isolated conditioned media from a Plant Cell Culture.

“Plant Cell Extract” is an extract of the cells (without the media).

Callus cells which are caused to differentiate into a specific plant part after callus formation are named by the plant part rather than callus term.

When the material is derived from a microorganism culture (with no additive), the name will consist of the genus and/or species term followed by the relevant term to indicate post-fermentation processing, (e.g., Lactobacillus Ferment Filtrate.)

Products derived by spontaneous fermentation (i.e., where a microorganism is not utilized) are named based on the material being fermented followed by the relevant term to indicate post-fermentation processing, (e.g., Sapindus Mukorossi Fruit Ferment Extract). The definitions for these ingredients indicate that the fermentation occurs spontaneously.

Materials produced through the cultivation of algae (or other microorganisms) are named on the basis of their composition and purity. The name is based on the identity of the species and the end-product, e.g., Mortierella Oil, Chlorella Protothecoides Oil, Euglena Gracilis Polysaccharide. See also 28 (a), and 43 (j).

Products derived by the fermentation of a microorganism which has been recombinantly modified are named by the prefix r- followed by the name of the microorganism.

Botanicals

Botanicals are cosmetic ingredients directly derived from plants. Generally, these ingredients have not undergone chemical modification and include extracts, juices, waters, distillates, powders, oils, waxes, saps, tars, gums, unsaponifiables, and resins. Where evidence of isolation is presented, a botanical ingredient may be named as a chemical entity, e.g., Genistein, or other appropriate terminology, e.g., Soy Isoflavones, depending on the extent of isolation.

The INCI names for botanicals are based on the Linné system or Latin binomial, whereby the Genus and species of the plant is used. In limited cases, the term related to the sub-species or variety has been historically used to differentiate materials that relate to the same Genus/species, e.g., Brassica oleracea captitata.

If an ingredient is derived from an interspecies botanical hybrid (same genus/different
species) with no recognized Linnean name, the INCI name will reflect the Linnean names of both plants used to create the hybrid separated by a slash, e.g., Rubus Fruticosus/Idaeus Extract. If an ingredient is derived from an intraspecies botanical hybrid (same genus/same species but different varieties), the genus/species of the phenotypic parent will be used, e.g., Helianthus Annuus (Sunflower) Seed Oil. In cases where an ingredient is derived from a cultivated hybrid in which the species name is unclear (e.g., the parentage of the species is from one or more hybrids), a cultivar name may be used in conjunction with the genus term, e.g., Phalaenopsis Charm Sun Big Red Robe Flower Extract.

c. Historically, the primary reference used by the INC to establish the Latin binomial names for botanicals was Penso, G., Index Plantarum Medicinalium Totius Mundi Eorumque Synonymorum, O.E.M.F. Milano (1983) - ISBN No. 88-7076-027-8.

d. Due to the dynamic nature of plant identification, the scientific nomenclature for plants is continually being updated. In order to provide accurate information, and minimize the impact of INCI name changes, the current accepted scientific plant names, where they differ from the INCI name, are described in the monograph definition. Notable botanical authorities are consulted in this effort, along with the various sources listed below:

- Index Fungorum, http://www.indexfungorum.org/
- International Plant Names Index, http://www.ipni.org/ipni/plantnamesearchpage.do
- The Plant List, http://www.theplantlist.org/
- United States Department of Agriculture Germplasm Resources Information Network (GRIN), http://www.ars-grin.gov/cgi-bin/npgs/html/tax_search.pl
- United States Department of Agriculture Plants Data Database, http://plants.usda.gov/java/nameSearch

e. INCI names for botanicals include the part(s) of the plant from which the material is derived. When more than one part is utilized, the plant parts are listed in alphabetical order, separated by a slash. When the material is derived from the entire plant, no part is specified and the material is defined as being from the whole plant.

f. The INCI names for plant extracts prepared by solvent extraction are assigned names that identify the name of the plant and the solvent. When the extraction solvent is carbon dioxide, carbon dioxide is not included in the INCI name since it evaporates. Additionally, solvents are not identified in the INCI name in cases where the solvent has been driven off and not present in the final preparation.

g. Essential oils prepared by a steam distillation process yields two distinct fractions, a water-insoluble fraction and a water-soluble fraction. The water-insoluble fraction contains the term oil in the name, e.g., Eucalyptus Globulus Leaf Oil. The water-soluble
fraction contains water in the name, e.g., Camellia Japonica Leaf Water. When an ingredient is prepared by adding water to a material prepared by solvent extraction, the ingredient is named as a mixture, e.g., Water (and) Juniperus Communis Fruit Extract. The term “water” is typically utilized for materials that are derived from plants; although “water” may be used to name non-botanical materials that are produced by steam distillation, e.g., Caviar Water, Royal Jelly Water.

h. The term “powder” is applied to the names for botanical materials that have been mechanically ground, irrespective of particle size. The term “meal” and “flour” are commonly recognized consumer terms, and are utilized accordingly in names such as Corn Cob Meal, Soybean Flour, etc.

i. Where several botanical materials are combined before processing, e.g., extraction or distillation, the ingredient is named by the genus, species and part of each plant separated by a slash followed by the preparation term, e.g., Aesculus Hippocastanum Bark/ Daucus Carota Root/Foeniculum Vulgare Fruit Extract. Exceptions to this convention are the INCI names Rose Extract and Rose Flower Oil which have historical usage and are grandfathered; along with Camellia Seed Oil defined as the oil expressed from one or more species of Camellia and named in accordance with information obtained from JCIA.

j. The term “soybean” has historical usage in INCI names to describe both Glycine soja and Glycine max.

k. Botanicals are named by a specific component, e.g., Apigenin, Isoquercetin, when the material has been isolated, purified and chemically characterized. General nomenclature for botanicals (e.g., Soy Isoflavones, Hydrolyzed Ginseng Saponin, Cassia Angustifolia Seed Polysaccharide) is utilized to name materials in accordance with the extent of their purification. See 54d for Conventions related to optical isomers.

l. Gums are polysaccharides of natural origin found in woody elements of plants or seed coatings, and in various seaweeds. Gums of natural origin are designated by common name that identifies the source, e.g., Acacia Senegal Gum, Ghatti Gum, Natto Gum. Common names for gums derived by fermentation include Gellan Gum, Xanthan Gum. Ingredients derived by reaction with a gum generally do not include the term “gum” in the INCI name, e.g., Hydroxypropyl Guar.

m. Ingredients derived from plant tissue culture are named in accordance with their process; the naming principles are fully described in Biotechnological Materials and Ferments.

n. The term “defatted” is used for ingredients which have been treated to remove lipid material which is typically accomplished through solvent extraction. For example, Defatted Hydrangea Macrophylla Flower refers to a plant extract in which the final preparation is the remaining plant material as opposed to the extract.

o. Seedcake is the term applied to the residue obtained after the oil has been processed from the seed.

p. Resins are secreted by specialized plant structures and consist of a mixture of terpenoid and phenolic compounds. Resins are often referred to as a sap, balsam, or exudate.
30. Harmonized INCI names for botanicals are designated by the Latin binomial as determined above, followed by the common name (where historically used) in parentheses, followed by the plant part (if applicable) and the type of preparation, e.g., Prunus Persica (Peach) Leaf Extract. See the Labeling Reminders section for an explanation of usage of common plant names.

In general, Latin binomial names are not used for botanicals that have been chemically modified. Botanicals that have a widely recognized common name (e.g., Olive Oil), and have undergone chemical modification may be named by common name and type of process, e.g., Acetylated Castor Oil, Hydrogenated Rapeseed Glycerides, Hydrolyzed Corn Starch, Oxidized Hazel Seed Oil, Ozonized Olive Oil, Saccharomyces/Grape Ferment Extract. In the absence of a previously monographed common name, or common name not widely known, the genus/species name may be utilized to name derivatives, e.g., Schinziophyton Rautanenii Oil Polyglyceryl-6 Esters.

In the EU, botanicals are named by the Latin binomial as explained above, followed by the plant part (if applicable) and type of preparation, e.g., Prunus Persica Leaf Extract. (See the discussions on international harmonization and botanicals in the Labeling Reminders section.)

When several materials relate to the same genus/species are used, the variety or sub-species in the Linné system may be identified, e.g., Citrus aurantium dulcis, Citrus aurantium amara.

There are a few cosmetic ingredients of herbal medicine origin in Japan in which the medicinal effect of the plant is specific to the sub-species. For these ingredients, the sub-species is included in the name, e.g., Coix Lacryma-Jobi Ma-Yuen Seed, Coix Lacryma-Jobi Ma-Yuen Seed Oil.

31. The INCI names for extracts represent the “material extracted”. The extracting solvent(s) if present in the final preparation is included in the INCI name assignment in descending order of concentration.

**Ceramides**

32. The term ceramide as part of an INCI is assigned to those classes and structures of natural lipids derived from skin as reported by Philip W. Wertz, Ph.D., Marion C. Miethke, M.D., Sherri A. Long, M.D., John M. Strauss. M.D., and Donald T. Downing, Ph.D., “The composition of ceramides from human stratum corneum and from comedones,” *The Journal of Investigative Dermatology*, 84 410-412 (1985). The term “ceramide” is also utilized in accordance with the naming system proposed by Motta, S., et al (1993) Biochimica et Biophysica Acta, 1182, 147-151.

a. A synthetic N-acylated sphingoid base that is identical to any one of the many constituents of the natural ceramides as reported by Wertz, has historically been assigned an INCI labeling name in accordance with the Wertz system, e.g., Ceramide 1, Ceramide 1A, Ceramide 2, Ceramide 3, Ceramide 4, Ceramide 5, Ceramide 6II. The term
ceramide as part of the INCI name will be assigned to a N-acylated sphingoid base that contains, as the predominant component, the D-erythro isomer of at least one of the many natural ceramides described by Wertz. A predominant component is one that is present at the highest concentration in relation to other synthetic materials of similar structure and related compositions present in a mixture. The Motta system for naming ceramides is also incorporated into INCI nomenclature. The Motta system utilizes a series of acronyms to designate the various fatty acid/sphingoid base combinations. The sphingoid base is typically 6-hydroxysphingosine, phytosphingosine, sphinganine or sphingosine, and the fatty acid can be saturated or unsaturated, and normal, or contain an alpha- or omega-hydroxyl grouping. Ceramides containing an omega-hydroxy fatty acid can exist in the free form or be esterified with either linoleic acid or a mixture of predominantly linoleic acid in combination with oleic acid and stearic acid. A number of different combinations of fatty acid/sphingoid base exist which give rise to a variety of INCI names, e.g., Ceramide NS, Ceramide EOS, etc. The chart below identifies the acronyms used in Motta-based ceramide nomenclature:

b. Synthetic N-acylated sphingoid bases that do not have the D-erythro configuration, or otherwise are not constituents of natural ceramides as described by Wertz or Motta, will not be named using the term ceramide. In such cases, a chemical, or other appropriate name, to be determined by the International Nomenclature Committee (INC) on a case-by-case basis, will be assigned as the INCI labeling name.

**Color Additives**

33. Color additives permitted for products to be marketed in the United States are identified in Title 21 of the *U.S. Code of Federal Regulations* (21 CFR). The INCI Names for color additives subject to batch certification are abbreviated names as identified in the *Federal Register* on June 6, 1985 (50 FR 23815). The abbreviated labeling names do not include “FD&C” or “D&C,” “No.,” or the type of lake “Aluminum, Zirconium, etc.,” on their product labels, e.g., Blue 1 Lake is the INCI name for the batch certified colorant FD&C Blue No. 1 Aluminum Lake.

b. For U.S. FDA batch certified colorants, additional names have been added as synonyms in order to identify the non-certified commodity, e.g., Pigment Red 57 instead of Red 7.

c. Alternative Color Index (CI) names have been established for those color additives appearing in Annex IV of *Regulation (EC) No 1223/2009 on cosmetic products* and are required to be used on products labeled for the European Union.
d. Alternate INCI names have been established for synthetic organic color additives permitted in Japan, regulated by the Ordinance to Regulate Coal-Tar Colors Permitted for Use in Drugs, Quasi-drugs, and Cosmetics (MHLW Ordinance No. 30 of August 31, 1966 as amended by MHLW Ordinance No. 55 of December 13, 1972, by MHLW Ordinance No.1126 of July 29, 2003 and by MHLW Ordinance No. 59 of May 2004

e. Coated pigments are named as blends, e.g., Polyethylene Terephthalate (and) Aluminum Powder. Epoxy Resin Coated Aluminum Powder is contained in the Dictionary as one of the Japan Trivial Names.

Oxidative hair coloring intermediates are named as described in 21 CFR. Those intermediates not appearing in 21 CFR are named according to their chemical structure.

Preformed hair colors are named as described in 21 CFR. Those preformed hair colors not appearing in 21 CFR are given the Colour Index Name. Preformed hair colors not appearing in either 21 CFR or the Colour Index are assigned chemical names based on their structure. In the event that the chemical name is very complex, these colors are assigned an arbitrary color/number designation, prefixed by the letters “HC.”

**Denatured Alcohol**

36. a. Specially Denatured (SD) Alcohols used in products marketed in the United States are named in compliance with Title 27 of the *U.S. Code of Federal Regulations* (27 CFR). The denaturants used in the manufacture of each SD Alcohol formula are specified in the monograph in Section 1. Manufacturers using these SD Alcohols should consult 27 CFR and the *Federal Register* for permitted uses, restrictions, and proposed changes.

b. An alternate INCI name, Alcohol Denat., has been established for products marketed in European Union (EU) Member States. Alcohol Denat. is ethyl alcohol that is denatured in accordance with the national legislation of each EU Member State. The INCI Name Alcohol Denat. may also be used in the United States for ethyl alcohol denatured in accordance with 27 CFR. For additional information see “Regulatory and Ingredient Use Information,” Introduction, Part A.

**Glycerides**

37. a. The term “Glyceride” has been utilized to describe a monoglyceride. (e.g., Acetylated Lard Glyceride, Canola Oil Glyceride, C10-40 Isoalkyl Acid Glyceride, Palm Glyceride.)

b. The term “Glycerides” is used to designate mixtures of mono-, di- and/or triglycerides, (e.g., Acetylated Palm Kernel Glycerides, Caprylic/Capric Glycerides, Corn Glycerides, Isostearic/Myristic Glycerides, PEG-12 Palm Kernel Glycerides.)

c. Triglycerides are designated by the term “triglyceride”. Alternate nomenclature is utilized when triglycerides are formed utilizing a single fatty acid, (e.g., Trilaurin, Trimyristin, Tristearin.)

**Glycols**

38. a. Glycol is the INCI name for ethylene glycol and is used as a combining term for derivatives of ethylene glycol, e.g., Glycol Distearate, Glycol Salicylate.
Alkylene 1,2-diols are usually named by the common name of the alkyl group followed by the term glycol, e.g., Lauryl Glycol. One exception includes hexylene glycol. See Convention 44f for the naming of polyethylene glycol.

b. Diglycol is the INCI name for diethylene glycol, and is used as a combining term for derivatives of diethylene glycol, e.g., Ethoxydiglycol, Diglycol/Isophthalates/SIP Copolymer.
   i. Diglycol is typically used in polymer names where diethylene glycol is used as a starting monomer, e.g., Bis-HEMA Poly(Diglycol Adipate)/IPDI Copolymer.
   ii. Diglycol is typically used in ethers, e.g., Butoxydiglycol, Methoxydiglycol.
   iii. Diethylene Glycol is typically used in esters, e.g., Diethylene Glycol Benzoate, Diethylene Glycol Dimethacrylate

c. PEG-2 is the INCI name for diethylene glycol when the reaction mechanism occurs through the use of an average of 2 moles of ethylene oxide. The exception to this principle is the usage of the suffix “-eth-2” to describe a 2-mole ethoxylate of a fatty alcohol, (see Convention 18a).

d. Butylene glycol is the INCI name for 1,3-butanediol. The numbers are omitted from the INCI name for the parent compound and its derivative, e.g., Butylene Glycol Myristate. INCI names for all other configurations include the numerical prefix to specify the position of the hydroxyl groups, e.g., 1,2-Butanediol, 2,3-Butanediol, 1,4-Butanediol, PEG/Poly(1,2-Butanediol)-52/32 Dimethyl Ether, 1,4-Butanediol Bisdecanoate. See also Convention 4.

e. Propylene Glycol is used as a combining term in INCI names when it is a starting material, e.g., Propylene Glycol Behenate, Dipropylene Glycol Caprylate, Tripropylene Glycol Citrate.
   i. “-PG” or “PG-” is used in INCI names when the starting material is glycidol, or where a material is reacted with one mole of propylene oxide.
   ii. Hydroxypropyl refers to 2-hydroxypropyl or 3-hydroxypropyl (e.g., Guar Hydroxypropyltrimonium Chloride), and is used when the starting materials are possibly 1-chloroisopropanol or 1-aminoisopropanol, or 3-chloropropanol or 3-aminopropanol. (See also Convention 19b)

**Hydrocarbons**

39. a. Hydrocarbon mixtures (notably solvents) not named by source or predominant feedstock as described in Conventions 20, 21, 22 are named on the basis of 80% of the composition utilizing a carbon number prefix which defines the 80% range as follows:
   i. Both numbers of the range will be even for materials containing only even-numbered carbon chains
   ii. Both numbers of the range will be odd for materials containing only odd-numbered carbon chains
iii. One number of the range will be even and the other odd for materials containing both even- and odd-numbered chains

b. A mixture of linear saturated hydrocarbons is named “Alkane” e.g., C14-22 Alkane. Ethoxylated derivatives of linear saturated alcohols are named in the same manner utilizing the term “Alketh”, e.g., C12-15 Alketh-3.

c. A mixture of branched saturated hydrocarbons is named “Isoalkane”, e.g., C32-54 Isoalkane, where the mixture can contain one or more branched substituents.

d. Cyclic saturated hydrocarbons are named “Cycloalkane”.

e. Mixtures of structurally different saturated hydrocarbons are named by each component separated by a slash, e.g., C9-16 Alkane/Cycloalkane.

f. Mixtures of unsaturated hydrocarbons are named “Olefin” and include mono- and polyunsaturates, linear, branched, and cyclic, e.g., C18-26 Olefin, Poly(C4-12 Olefin).

Imidazolines

40. Common fatty stem terms are used to designate the alkyl portion of alkyl imidazoline compounds (e.g., Lauryl Hydroxyethyl Imidazoline) even though one carbon atom of the fatty radical becomes a member of the heterocyclic ring during the materials’ manufacture.

Lanolin Derivatives

41. Names of lanolin derivatives usually contain the stem “lan”, e.g., Laneth-10 Acetate. When fractionated, derivatives are named utilizing “lan” as a stem name unless a specific component has been isolated, e.g., Cholesterol.

Minerals

42 a. Naturally occurring minerals with a definite chemical composition and/or physical properties (which may include x-ray diffraction data) are named according to the established, published nomenclature. Some reference sources include:

- www.mindat.org
- http://minerals.usgs.gov/

b. Naturally occurring materials that are mixtures of mineral species are named on the basis of particle size using common names such as sand, clay, silt, and other similar terms. Historically, some inorganic materials have been named according to geographic origin when the composition and properties with regard to origin were properly documented and supported in the literature, e.g., Moroccan Lava Clay.

c. The term “synthetic” is applied to the names of inorganic materials such as rocks, gems, and minerals, (e.g., Synthetic Ruby) to indicate that the material is synthesized. These materials, while generally physically indistinguishable from their natural counterparts, are chemically similar but may vary in chemical composition. Bureau of Standards and X-ray diffraction pattern data must be supplied to support the characterization of
compositional similarities between natural and synthetic materials.

d. Rocks, gems, and minerals that are mechanically ground (i.e., not ground by natural processes) are named by the common geological term followed by the term “powder”, (e.g., Ruby Powder).

e. Doped minerals obtained via calcination are considered solid solutions and are named as a single entity by the constituent mineral oxides, e.g., Silicon/ Titanium/Cerium/Zinc Oxides.

f. Mineral extracts are designated by the name of the mineral and the term “extract”, e.g., Loess Extract, Lignite Extract, Malachite Extract, when the manufacturing information indicates the mineral is extracted.

g. Allotropes of carbon are named according to their structural form, e.g., Diamond, Graphite, Fullerenes, Carbon.

h. Plant Ash is the name designated for ingredients composed of ash produced by the combustion of any plant material or mixture of plants.

i. Carbonaceous material obtained by heating wood or other organic matter in the absence of oxygen is named as Charcoal.

j. Clay as an INCI name refers to a group of phyllosilicate minerals produced by the chemical and physical weathering of rock. It consists chiefly of varying amounts of hydrated silica and alumina and is characterized by a particle size of less than 2 micrometers. Clays have historically been designated INCI names based on geographic region; these names have been “grandfathered”, e.g., Heilmoor Clay. The current naming practice is to designate clay materials by the INCI name, “Clay”. See also 42 (b).

k. Loose pieces of minerals and rocks are sediment and further characterized by particle size as follows:
   i. Sand is a naturally occurring granular material composed of finely divided rock and mineral particles based on silica in the form of quartz, with a typical particle size between 0.0625 to 2.00 mm.
   ii. Silt is sediment from inland bodies of water. It is a naturally occurring inorganic material whose origins are based on quartz and feldspar with typical particle sizes between 0.0625 to 0.00400 mm, e.g., Sea Silt.
   iii. Mud is a mixture of water and some combination of soil, silt, and clay, e.g., Alluvial Mud, Salt Mine Mud.

l. Volcanic Soil is a mixture of minerals derived from volcanic deposits which are of varying size including but not limited to sand, silt and clay. Volcanic Sand is loose, granular particles of disintegrated lava deposits. Volcanic ash is the residue obtained from volcanic eruption.

m. Glass is an amorphous inorganic material based on silica (SiO2) that is combined with various additives, usually metal oxides (e.g., sodium oxide, calcium oxide, magnesium
oxide). Glasses are produced by fusing silica together with the additives, then rapidly cooling to eliminate formation of a crystalline structure.

n. INCI names do not generally include descriptive terms related to particle size. Materials that result from milling are named as powders or flours irrespective of size. See Convention 67 for usage of the term colloidal. Additionally, “nano” is not used in the assignment of INCI names to trade materials. See Convention 73.

Oils, Fats, Lipids and Triglycerides

43. a. Triglycerides of plant or animal origin that are liquid at room temperature are generally known as fixed oils and are named by their source followed by the term oil, e.g., Olea Europea (Olive) Oil, Elaeis Guineensis (Palm) Oil, Elaeis Guineensis (Palm) Kernel Oil, Canola Oil, Cod Liver Oil. Oils which have been chemically modified by acetylation, hydrogenation, isomerization or oxidation are named in a similar fashion, e.g., Acetylated Castor Oil, Hydrogenated Palm Kernel Oil, Isomerized Palm Oil, Oxidized Corn Oil.

b. Triglycerides of animal origin that are solid at room temperature are generally known as fats and are named by their source followed by the term fat e.g., Deer Fat, Goat Fat, Buffalo Fat.

c. Essential oils that are water insoluble fractions of plant materials obtained by steam distillation are named by their source followed by the term oil, e.g., Rose Flower Oil, Salvia Officinalis (Sage) Oil. Water soluble fractions of essential oils are named as waters.

d. The term “oil” may be used to name non-triglycerides when it applies to ingredients that are commonly recognized, (e.g., Simmonisia Chinensis (Jojoba) Oil, Lanolin Oil, Mineral Oil, Tall Oil, Tar Oil.)

e. Plant butters derived by mechanically pressing the seeds are generally semi-solids at room temperature and are named by the genus/species term of the plant in accordance with Convention 30, e.g., Garcinia Indica Seed Butter. Exceptions include the common name “Butter” which refers to the fat recovered from cow’s milk, and “Goat Butter” which refers to the fat recovered from goat’s milk.

f. Lipids isolated from plant or animal origin are named by the common name of the animal, e.g., Shark Lipids, Silkworm Lipids, or genus/species name of the plant, e.g., Oryza Sativa (Rice) Lipids. See also Convention 33, Ceramides.

g. Lipids produced by various strains of algae are named in accordance with the composition of the final product. If the product is constituted by a mixture of fatty acids, approximately 80% of the fatty composition is identified in alphabetical order separated by a slash, e.g., Capric/Lauric/Myristic/Oleic Triglyceride.

Polymers

44. Due to the wide variety of polymer chemistry types exhibited among synthetic and naturally occurring polymers and due to their inherent polydispersities, there is no uniform approach to distinguish between what constitutes an oligomer vs. a polymer
solely on the basis of degree of polymerization (i.e. the number of repeat units in the polymer chain). Instead, the oligomer vs. polymer distinction on the basis of degree of polymerization is specific to individual polymer chemistry type, and it is typically associated with measurable differences in macroscopic physical properties, such as melting point, viscosity, etc., that occur as a function of degree of polymerization for that polymer chemistry type.

Polymeric materials are named according to the name in common usage if it is well known, or by the structure if well-defined, e.g., polyethylene terephthalate. Typically, polymers are named by the starting monomer instead of the composition of the final polymer, e.g., Polydecene instead of polydecane. Exceptions include copolymers named with vinyl alcohol as one of the monomers, e.g., Sodium MA/Vinyl Alcohol Copolymer, in which the starting monomer, vinyl acetate is hydrolyzed to form the alcohol. If no common name exists, and the structure is not well-defined, the polymers are named according to their composition as described below:

a. Homopolymers (consisting of one constituent monomer) are named by placing the term “poly” before the constituent monomer, e.g., Polyisobutene.

b. Copolymers consisting of two or more constituent monomers are named by listing the monomers in alphabetical order separated by a slash (/) followed by the word “Copolymer,” e.g., Acrylates/Acrylamide Copolymer.

c. Copolymers consisting of four or more monomers may be given an INCI name according to the predominant monomer, or resultant polymer class followed by an arbitrary number, e.g., Polyester-1, Polyquaternium-1, etc., with the monomers listed in the monograph definition in alphabetical order of the material. Copolymers with less than four monomers and with an excessive name length may also be considered for “polytype” names as described above. Such nomenclature is granted at the discretion of the INC.

d. Crosspolymers consisting of two or more constituent monomers are named by listing the monomers in alphabetical order separated by a slash (/) followed by the word “Crosspolymer,” e.g., Acrylates/VA Crosspolymer.

i. The crosslinking agent will be included in the INCI name if the crosslinking agent is a polymer. In these cases, the crosslinking agent will appear as the last component of the INCI name followed by the word “Crosspolymer”; e.g., Lauryl Dimethicone/PEG-15 Crosspolymer, where the crosslinker is diallyl PEG-15. When the crosslinking agent is not a polymer, it will not be included in the INCI name, but will be included in the monograph definition of the material, e.g., Acrylic Acid/Isopropylacrylamide/ MIBK Acrylamide Crosspolymer-is a copolymer of acrylic acid, isopropylacrylamide, methyl isobutyl ketone (MIBK) acrylamide crosslinked with methylene bis-propenamide. Carbomer is an exception to this Convention because of its historic usage.

ii. In cases where a polymer cannot be formed in the absence of a crosslinking monomer, i.e., the crosslinking monomer is essential for the formation of the polymer repeating structure, the crosslinking monomer will be included in the
INCI Name. For example, the crosslinked polyester formed by the condensation of Propanediol and Citric Acid would be named Propanediol/Citrate Crosspolymer. Exceptions include polymers named in accordance with Convention 44(c).

45. The term “Acrylates” is used to describe linear, non-crosslinked copolymers that contain combinations of acrylic acid, methacrylic acid, and their simple esters. They are described as simple alkyls ranging from C1 to C4 (linear or branched). Similarly, the term “Crotonates” is used to describe copolymers that contain combinations of crotonic acid and its simple esters.

46. The term “Aminoacrylates” refers to simple aminoacrylates, in which the substituted alkyl groups attached to amino nitrogen range from C1-4, and acrylates conforms to the definition as described above.

47. The name “Carbomer” is used to describe high molecular weight crosslinked homopolymers of acrylic acid. The crosslinking agent(s) are identified in the ingredient monograph definition.

48. A “Dendrimer” polymer is named from the core to the outside by the monomer layers. If a monomer unit is repeated, the number of generations or layers is indicated. If a previous convention exists for naming the core, then it is utilized. An example is PEG-5 Pentaerythrityl (the core) Dimethylol (the layer) Propionate-2 (generations) in which there are 5 repeating units of polyethylene glycol attached to pentaerythritol as the core. Dimethylol propionic acid is reacted to the core for two generations.

A dendron attached to a polymer backbone is named by the backbone polymer with the added dendron side group described, e.g., Acrylates/HEMA Copolymer (the core) Dimethylol Propionate-4 (the layer and generation) Dendron.

49. The term Polyurea is used to name polymers typically formed by the condensation of a diisocyanate with a diamine.

Prostaglandin Derivatives

50. Ingredients which are analogues of prostaglandin compounds utilize the drug stem term as part of the corresponding cosmetic ingredient name, e.g., Bimatoprost and Cyclopentylbimatoprost; Cloprostenol and Isopropyl Cloprostenate; Noralfaprostol and Isopropyl Dihydro Noralfaprostol; Travoprost and Ethyl Travoprostamide. The use of common drug stem names for related cosmetic substances is considered by the INC on a case-by-case basis.

Quaternary Ammonium Salts

51. Quaternary ammonium salts usually have the suffix “ium” in the stem of the cation. The term “monium” describes a monomethyl-substituted quaternary nitrogen; “dimonium” describes a dimethyl-substituted quaternary nitrogen; “trimonium” describes a trimethyl-substituted quaternary nitrogen.

Silanes and Siloxanes

52. Silanes and Siloxanes are named according to the following subcategories:
a. Silanes are monomeric compounds containing one silicon atom. They are used as intermediates to prepare siloxanes and to modify the surfaces of other ingredients. Silanes are named by listing substituents in alphabetical order, and then the term ‘silane’ with the appropriate numerical prefix (e.g., Trimethoxycaprylylsilane).

b. Silanols [silanes containing hydroxyl group(s)] are named according to the number of hydroxyl groups attached to the silicon atom. The other substituents included in the name (e.g., Hydrolyzed Keratin PG-Propyl Methylsilanediol, Silanetriol Lysinate).

c. Siloxanes, commonly referred to as silicones, are polymers that are based on chains of alternating silicone and oxygen atoms that also contain organic substituents. The most common organic substituents are methyl (-CH₃) but many other types of organic substituents are possible. The most common siloxanes are linear polysiloxanes with two methyl substituents on each silicon atom (polydimethylsiloxanes). When the polydimethylsiloxanes are terminated with methyl groups, the INCI name is Dimethicone. Short chain Dimethicones (2-4 silicon atoms) are named chemically (e.g., Trisiloxane, Disiloxane). Dimethicones where some of the methyl groups along the chain are replaced with other organic groups are named as Dimethicone derivatives (e.g., Stearyl Dimethicone, PEG-8 Dimethicone). It is assumed the substituents on Dimethicone derivative are attached through a propyl linkage. Any other attachment is identified in the name.

d. Dimethicones where the polysiloxane chain is terminated with substituents other than methyl groups are named using “Bis-” to indicate where the substituents are located (e.g., Bis-PEG-8 Dimethicone, Bis-Hydroxypropyl Dimethicone)

e. Dimethicones where the polysiloxane chain is terminated with hydroxyl groups (-OH) are called Dimethiconols. Hydroxyl groups attached to silicon atoms are somewhat reactive. They can condense with each other to form a new siloxane bond (-Si-O-Si-), leading to chain extension. They can also react with organic acid to form a type of ester (e.g., Dimethiconol Behenate)

f. Methicone refers to linear siloxane polymers where each silicon atom in the siloxane chain has one methyl group and one hydrogen atom. Methicone is a reactive polymer that is used to create a hydrophobic coating on the surface of inorganic pigments. It is also the starting material for making alkyl siloxanes where the hydrogen atoms are replaced with the alkyl group (e.g., C26-28 Alkyl Methicone).

g. Cyclic dimethyl siloxane was historically named ‘Cyclomethicone’ to represent mixtures of species containing three to seven siloxane units. For pure components (>99%), the nomenclature is based upon the number of siloxane units: Cyclotrisiloxane, Cyclo-tetrasiloxane, Cyclopentasiloxane, and Cycloheptasiloxane. In the silicone industry, these cyclic dimethyl siloxanes are often referred with the following shorthand notation: D3, D4, D5, and D6.

h. Silsesquioxanes are highly branched siloxanes where each silicon atom is connected to three oxygen atoms and conform to the general formula [RSiO_{3/2}]x, where R is an organic substituent. They are sometimes referred to as Polyhedral Oligomeric Silsesquioxanes (POSS). The nomenclature is understood to include oligomeric
silsesquioxanes that only contain 4-6 silsesquioxane units (cage structures). If the organic group for the silsesquioxide has a short, simple name then the INCI will be a single word, e.g., Polyphenylsilsesquioxide, Polypropylsilsesquioxide. For more complex organic groups, they will be used with the term Polysilsesquioxide (e.g., Acryloyloxypropyl Polysilsesquioxide, Glycidoxypropyl Polysilsesquioxide). Silsesquioxanes frequently contain hydroxyl groups attached to the silicon atoms that allow these silsesquioxanes to react with other types of siloxane polymers. These are named as copolymers (e.g., Dimethicone/Silsesquioxane Copolymer).

i. Silicates are inorganic polymers where each silicon atom is attached to four oxygen atoms and conform to the general formula \([\text{SiO}_4^{4/2}]x\). If no organic substituents are present, then the INCI name is Silicon Dioxide. Silicones that contain silicate units are named as such, with any substituents and/or terminal groups appropriately named (e.g., Trimethylsiloxy silicate).

j. The term ‘Polysilicone’ followed by an arbitrary number is used to describe complex silicone polymers that cannot be named by common names or established conventions for silicone compounds (e.g., Polysilicone-10).

**Substituted Compounds and Prefix/Suffix Terms**

53. Singly substituted derivatives usually do not include the prefix “mono.” This term is used only when required to prevent ambiguity. The absence of a suitable prefix implies “mono,” e.g., Glyceryl Stearate represents glyceryl monostearate, and Glyceryl Oleate/Laurate represents a monoester of glycerin with a blend of oleic and lauric acids.)

54. Multiple substitution is routinely described with the appropriate prefix, such as “di-,” “tri-,” or “tetra-,” e.g., Glyceryl Distearate, Propylene Glycol Dilaurate, Pentaerythrityl Tetra benzoate.

a. Where there is substitution with a mixture of components, i.e., alkyl groups, the prefix is used only once, wherever possible, and the moieties are separated by a slash, e.g., Ditrimethylolpropane Tetraisostearate/ Hydroxystearate to denote the tetraester of ditrimethylolpropane and a mixture of isostearic and hydroxystearic acids.

b. The simple numerical prefixes “di-”, “tri-”, “tetra-” etc. are used to indicate a multiplicity of simple (i.e., unsubstituted) substituents provided that there is no ambiguity, e.g., Propylene Glycol Dilaurate, Triethyl Citrate.

c. The numerical prefixes “bis”-, “tris”- are generally utilized to denote multiple identical structural features of a compound, e.g., Tris-Biphenyl Triazine, Bis-Aminopropyl Dimethicone.

d. Optical isomers are usually not designated in INCI names although this information may be included in the monograph definition. However, there may be circumstances whereby it is necessary to identify the optical properties of the isomer, e.g., d-limonene under EU regulation 111/1.88, and the INC will address these situations as they arise.
e. The numbering of substituents is only employed where necessary to prevent ambiguity, e.g., 1,4-Butanediol, 2,3-Butanediol.

f. The prefixes o-, m-, p-, t-, n-, N, N', etc. are used only when necessary to prevent ambiguity.

g. Locants are included in the INCI name when there is more than one possible site for the reaction. e.g., N-Feruloyl Dopamine

55. Mixtures of mono-, di- and tri-esters of glycerin are designated by the suffix “-ates”, (e.g., Glyceryl Stearates.)

56. The dimethyl term is omitted and is assumed in all alkyl dimethyl amine oxide names (e.g., Stearamine Oxide). Tertiary amine oxides with different substituent groups are named completely (e.g., Dihydroxyethyl Stearamine Oxide).

57. Amino acids substituted on nitrogen are named by the identity of the substituent group and the trivial name of the amino acid. Since N-2 is the atom most easily modified, N-can be omitted from the name without ambiguity, e.g., Acetyl Tyrosine.

a. The suffix “ate” is added to the amino acid name when the substance is a salt, e.g., Sodium Glutamate, Potassium Aspartate, or an ester, e.g., Ethyl Glutamate, Acetylated Cetyl Hydroxyproline, Methyl Undecenoyl Leucinate.

b. When the hydroxyl group of the carboxyl has been replaced by an amino group, the “amide” suffix is added to the trivial name of the amino acid, e.g., Hexacarboxymethyl Lysinyl Lysinamide, Hydroxyphe nylic Glycinamide, Prolinamidoethyl Imidazole.

c. When amino acids are derivatized, amino acid stem names are used as a combining term rather than chemical names, e.g., Glutamyl Hydroxyphenylhydrazide, Prolyl Histidine HCl, Palmitoyl Lysyl Aminovaleryl Lysine.

58. The prefix “dimer” precedes the term “dilinoleic” to designate materials that are C36 di-acids; it has historical usage in INCI nomenclature, e.g., Dicetearyl Dimer Dilinoleate.

59. The prefix “nor” is used to designate “de-methyl” which means one methyl group removed relative to the parent compound for the purposes of nomenclature, e.g., Norvaline.

60. The prefix sesqui- is used for esters of alcohols and fatty acids that are mixtures of the monoester and diester, e.g., Butylene Glycol Sesquisostearate, Methyl Glucose Sesquilaurate, PEG-8 Sesquioleate, Polyglyceryl-2 Sesquicaprylate.

**Synthetic and Recombinant Peptides**

61. Natural protein ingredients, and their derivatives, are named by the common name of the source, e.g., Albumen, Collagen, Fibroin, Mellitin, Milk Protein, Nacre Protein, Serum Protein, Silk, etc. See also 28h. Peptides, produced either by chemical synthesis or recombinant techniques, are named based on sequence and by the following rules:
a. Synthetic peptides consisting of two to ten amino acid residues are named using the appropriate prefix, di-, tri-, tetra-, etc., followed by the term peptide and an arbitrary number, e.g., Dipeptide-2, Decapeptide-4, Pentapeptide-3. The constituent amino acids are identified in the monograph definition. There are a few peptides that are historically named by their amino acids, e.g., Glycyl Glycine, and these names have been grandfathered. Additionally, Glutathione is a grandfathered name for the peptide, glutamyl cysteinyl glycine, whereas Tripeptide-35 is composed of the same amino acids but of possible differing sequence.

b. Synthetic peptides consisting of 11 to 100 amino acids are designated by the term oligopeptide, followed by an arbitrary number, and the constituent amino acids are identified in the monograph definition, e.g., Oligopeptide-13.

c. Synthetic peptides consisting of more than 100 amino acids are designated by the term polypeptide, followed by an arbitrary number, and the constituent amino acids are identified in the monograph definition, e.g., Polypeptide-5.

d. The amino acid residues composing the peptide are listed alphabetically in the monograph definition. The amino acid residues may include the following: Alanine, Arginine, Asparagine, Aspartic Acid, Cysteine, Glutamic Acid, Glutamine, Glycine, Histidine, Isoleucine, Leucine, Lysine, Methionine, Phenylalanine, Proline, Serine, Threonine, Trptophan, Tyrosine, Valine.

e. When the peptide contains an amino acid that is not one of the natural amino acids identified above, (e.g., D-isomers or gamma amino acids) it is identified in the peptide name, e.g., Tripeptide-9 Citrulline, Acetyl Norleucyl Dipeptide-54 D-Phenylalanyl Dipeptide-25 Amide, Histidyl D-Tryptophanyl Dipeptide-29 D-Phenylalanyl Lysinamide.

f. Peptide derivatives are named utilizing the parent peptide name, and the name of the modifying group as follows:
   i. When the N-terminus is modified, the name of the modifying group precedes the peptide name, e.g., Myristoyl Hexapeptide-5, Palmitoyl Octapeptide-24
   ii. When the C-terminus is modified, the name of the modifying group is identified after the peptide name according to its composition, e.g., Tripeptide-9 Citrulline, Caffeoyl Tetrapeptide-19 Caffeamide, Tetrapeptide-29 Argininamide, Acetyl Octapeptide-17 Amide, Tripeptide-83 Propyl Ester.
   iii. When any hydroxyl group or amine group along the peptide chain is modified, it is named according to the composition of the reacting species.

g. Ingredients originating from human tissue, including materials starting with human cells in tissue culture, contain “human” as part of the INCI name. See 27(b). See 61(h) for ingredients made by recombinant technology.

h. Names for biologically inspired peptides are based on the source protein whether produced through recombinant techniques or chemical synthesis. The name is constructed by prefixes to designate recombinant and synthetic, followed by the
peptide term as outlined in 61(a-f), followed by a number, whereby the number is unique and corresponds to the specific protein which is described in the ingredient definition.

i. The prefix “rh” is used to identify “recombinant human” peptides in which the original gene is isolated from a human cell, e.g., rh-Oligopeptide-1, rh-Polypeptide-, rh-Polypeptide-96.

ii. When the original gene is derived from another organism, the prefix “r-“ is used, followed by the name of the organism, e.g., r-Mussel Polypeptide-1, r-Bovine Octapeptide-1, r-Leucosporidium Polypeptide-1.

iii. When the gene is synthesized to be identical to a human gene and produced recombinantly, the prefix “sh” for “synthetic human” is used, e.g., sh-Polypeptide-96, sh-Oligopeptide-8.

iv. When the gene is a synthesized copy from another organism and the peptide is produced through recombinant technology, the prefix “sr-“ is used to indicate synthetic recombinant followed by the name of the organism, e.g., sr-Bovine Oligopeptide-2, sr-Jellyfish Polypeptide-1, sr-Rice Polypeptide-1.

v. Biologically inspired peptides that are produced by chemical synthesis are designated by the prefix “s-“, followed by the name of the organism, e.g., s-Centipede Decapeptide-1, s-Bovine Oligopeptide-1, s-Coconut Antimicrobial Peptide 1. When the species is human, the prefix “sh” is used and the suffix “SP” is added to the name to indicate solid phase synthesis, e.g., sh-Oligopeptide-23 SP, sh-Pentapeptide-6 SP. The latter approach was employed to distinguish these peptides from earlier ones produced recombinantly as described in 60(h)(iii). In some cases, the suffix SP is added to the peptide name to further distinguish the synthetic nature of the peptide, e.g., s-Mussel Hexapeptide-1 SP.

vi. Peptides that are derived from two or more different peptides are considered fusion proteins. They are named by combining the INCI names of each appropriate individual peptide name, e.g., r-Clostridium Histolyticum Collagenase sh-Oligopeptide-60; r-Mussel Polypeptide-1 r-Mussel Oligopeptide-1 sh-Polypeptide-1.

i. Antibodies are immunoglobulins (Ig), composed of 4 polypeptide chains (2 “heavy” and 2 “light” chains) bound to each other by disulfide bonds. The heavy chains are typically glycosylated. Antibodies are not named as “polypeptides” due to their complexity and unique structure. Rather than amino acid sequence, they are named according to antigen specificity.

Antibodies (immunoglobulins) are named according to the antigen they specifically bind, the organism where they are raised or the hybridoma cells they are isolated from, and their type, whether monoclonal (from hybridoma cell cultures) or polyclonal (from serum). The basic naming approach is as follows: Anti-Protein/Antigen Organism Type Antibody. The common name for the antigen is used when available. As an example, a monoclonal antibody raised against Collagenase-1 in murine hybridoma is named: Anti-Collagenase-1 Mouse
Monoclonal Antibody.

j. Proteins derived from a transgenic viable higher organism in which the gene expression is either stable or transient are named by the host organism, followed by the appropriate peptide name, and are defined by a transgenic process. Common names are often used to identify the host organism, e.g., Barley sh-Oligopeptide-1.

k. Peptides which maintain the full sequence of a protein and are sourced directly from an organism are identified by the name of the protein, e.g., Cobrotoxin, Alloferon-1.

l. Peptides which maintain the full sequence of a protein and are synthetically derived are identified by the name of the protein with an “s-“ prefix, e.g., s-Pentadiplandra Brazzeana Defensin-Like Protein, s-Mellitin, s-Mu-conotoxin CnIIIC.

m. Aptamers are synthetically produced single-stranded oligonucleotides that fold into a three-dimensional structure and bind to a specific target. They are named based on composition, e.g., DNA, RNA, followed by the term “aptamer” and a unique number, e.g., s-DNA Aptamer-2. The ingredient definition describes the number of constituent nucleotides and the target molecule to which it binds.

Transesters

62. Transesters are materials derived by the process of the transesterification of esters, usually triglycerides (fats and oils), and alcohols. They are generally identified by the term “esters”, e.g., Apricot Kernel PEG-8 Esters, when the alcohol is less than a stoichiometric amount, there is no purification, and the material consists of a complex mixture of products, including mono- and di-glycerides and alcohols. Transesters can also be obtained by the transesterification of an oil with another oil, e.g., Moringa Oil/Hydrogenated Moringa Oil Esters. If a stoichiometric amount or excess of alcohol is used, the material will be named using Ester nomenclature, e.g., Ethyhexyl Cocoate.

A similar process is carried out by the process of the transamidation of esters, usually triglycerides (fats and oils), and amines. They are generally identified by the term “amides”, e.g., Coconut Oil MIPA Amides, when the amine is less than a stoichiometric amount, there is no purification, and the material consists of a complex mixture of products, including mono- and di-glycerides and amides. If a stoichiometric amount or excess of amine is used and the reaction driven to completion, the material will be named using Amide nomenclature, e.g., Cocamide MIPA.

MISCELLANEOUS CONVENTIONS

63. Amidino is the root used to designate the structure R-C(NH2)=NH, e.g., Amidinoproline.

64. “Esylate” is the term used to designate ethanesulfonate, e.g., Sodium Esylate. “Tosylate” is the term used to designate toluenesulfonate, e.g., Cetrimonium Tosylate.

65. Estolides are esters formed by the polymerization of 2 or more hydroxyl fatty acids (e.g., 12-hydroxystearic acid), or by the acid-catalyzed condensation of 2 or more unsaturated fatty acids (e.g., oleic acid) to form oligomeric esters. The product is named “estolide” preceded
66. The compound commonly known as conjugated CLA is named as Isomerized Linoleic Acid.

67. Encapsulated materials are named as mixtures with the components identified in order of predominance.

68. Solutions that are characterized as the dispersion of very small particles in a continuous phase that remain suspended are named colloids, e.g., Colloidal Silver, Colloidal Platinum.

69. Zwitterions are internal salts and are identified as “betaines” and “sultaines”, e.g., Cocamidopropyl Betaine, Cocamidohydroxypropyl Sultaine.

70. Hydrocarbons derived by the complete hydrogenation of an unfractionated fatty acid are named according to the source of the fatty acids, e.g., Coconut Alkanes. See also Conventions 20 and 21.

71. “Lactylate” is the combining term used to describe the ester formed between two moles of lactic acid, e.g., Sodium Stearoyl Lactylate, Sodium Cocoyl Lactylate.

72. “Ascorbate” is used in INCI names for ingredients produced by the reaction of an alcohol with ascorbic acid to form an ether, e.g., Caprylyl 3-Glyceryl, or when a salt of ascorbic acid is formed, e.g., Calcium Ascorbate. “Ascorbyl” is used for esters between ascorbic acid and a fatty acid or phosphoric acid, e.g., Ascorbyl Stearate. Exceptions include the INCI names 2-O-Ethyl Ascorbic Acid, 3-O-Ethyl Ascorbic Acid, 3-O-Cetyl Ascorbic Acid.

73. Process terms are not usually utilized in INCI nomenclature, e.g., “heat-induced”. Exceptions include terms such as acetylated, epoxidized, extract, ferment, hydrolyzed, hydrogenated, lysate, ozonized.

74. For products marketed in the EU which contain ingredients that meet the definition for a nanomaterial as identified by the EC Regulation No. 1223/2009, the INCI name (Nano) may be applied as a suffix to the corresponding INCI name. For example, usage of titanium dioxide that meets the EC definition for nano would be declared on the finished product label as Titanium Dioxide (Nano).

75. Lactones are generally preceded by a Greek letter prefix to indicate the size of the lactone ring, e.g., Gamma- Caprolactone, Delta-Decalactone.

76. Glycosides are compounds which consist of a sugar (glycone) linked to a non-sugar or alcohol (aglycone) through a glycosidic bond formed through the condensation. The resulting compound is a named by the aglycone term first, followed by the sugar, where the “ose” suffix of the sugar name is replaced by “oside”. When the glycone is glucose, the resulting compound is named as a glucoside, e.g., Decyl Glucoside. Glucosides is used when the sugar is a polymer of glucose. When the glycone is xylose, the compound is named as a xyloside, e.g., Octyldecyl Xyloside, maltose would be a maltoside, etc.
When the aglycone is a complex material, such as a flavonoid or a ceramide, the sugar term precedes the aglycone, Glucosyl Hesperidin, Glucosyl Naringen, Glucosyl Ceramide NP, etc. Early exceptions to this convention include the names Phloridzinyl Glucoside, Polydatin Glucoside.

When the glycone is comprised of carbohydrates derived from a natural source, the material is named as a “glycoside”, preceded by the aglycone term, e.g., Cetearyl Wheat Bran Glycosides. Naturally occurring plant glycosides (e.g., saponins) may be named by their common or usual name, e.g., Ziyu Glycoside I.

**LIST OF ACRONYMS**

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<tr>
<td>AEEA</td>
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<td>Monoisopropanolamine</td>
</tr>
<tr>
<td>NTA</td>
<td>Nitrilotriacetic Acid</td>
</tr>
<tr>
<td>PABA</td>
<td>para-Aminobenzoic Acid</td>
</tr>
<tr>
<td>PCA</td>
<td>Pyrrolidone Carboxylic Acid</td>
</tr>
<tr>
<td>PEG</td>
<td>Polyethylene Glycol</td>
</tr>
<tr>
<td>PEI</td>
<td>Polyethylenimine</td>
</tr>
<tr>
<td>PG</td>
<td>Propylene Glycol</td>
</tr>
<tr>
<td>PPG</td>
<td>Polypropylene Glycol</td>
</tr>
<tr>
<td>PTFE</td>
<td>Polytetrafluoroethylene</td>
</tr>
<tr>
<td>PVM/MA</td>
<td>Polyvinyl Methyl Ether/Maleic Anhydride</td>
</tr>
<tr>
<td>PVP</td>
<td>Polyvinylpyrrolidone</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic Acid</td>
</tr>
<tr>
<td>SD</td>
<td>Specially Denatured</td>
</tr>
<tr>
<td>SE</td>
<td>Self-Emulsifying</td>
</tr>
<tr>
<td>SIP</td>
<td>Sulfoisophthalate</td>
</tr>
<tr>
<td>SMDI</td>
<td>Saturated Methylene Diphenyl-diisocyanate</td>
</tr>
<tr>
<td>TAED</td>
<td>Tetraacetylenediamine</td>
</tr>
<tr>
<td>TBHQ</td>
<td>tert-Butyl Hydroquinone</td>
</tr>
<tr>
<td>TDI</td>
<td>Toluene Diisocyanate</td>
</tr>
<tr>
<td>TEA</td>
<td>Triethanolamine</td>
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<tr>
<td>TIPA</td>
<td>Triisopropanolamine</td>
</tr>
<tr>
<td>TMHDI</td>
<td>Trimethylhexanediisocyanate</td>
</tr>
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<td>TMMG</td>
<td>Tetramethoxymethylglycouril</td>
</tr>
<tr>
<td>TMP</td>
<td>Trimethylolpropane</td>
</tr>
<tr>
<td>VA</td>
<td>Vinyl Acetate</td>
</tr>
<tr>
<td>VP</td>
<td>Vinyl Pyrrolidone</td>
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