Title: PREPARATION OF BRANCHED ALIPHATIC ALCOHOLS USING COMBINED PROCESS STREAMS FROM A HYDROGENATION UNIT AND A DEHYDROGENATION-ISOMERIZATION UNIT

Abstract: Systems and methods to produced branched aliphatic alcohols are described. Systems may include a hydrogenation unit, a dehydrogenation-isomerization unit, a hydroformylation unit and/or combinations thereof. Methods for producing branched aliphatic alcohols may include isomerization of olefins in a process stream. The isomerized olefins may be hydroformylated to produce aliphatic alcohols. After hydroformylation of the aliphatic alcohols, unreacted components from the hydroformylation process may be separated from the aliphatic alcohols products. The unreacted components from the hydroformylation process may be recycled back into the main process stream or sent to other processing units. Addition of multiple streams to the units may be performed to control reaction conditions in the units.
PREPARATION OF BRANCHED ALIPHATIC ALCOHOLS USING COMBINED PROCESS STREAMS FROM A HYDROGENATION UNIT AND A DEHYDROGENATION-ISOMERIZATION UNIT

Background of the Invention

Cross Reference to Related Application

This application claims the benefit of U.S. Provisional Application Serial No. 60/511,562 filed October 15, 2003.

Field of Invention

The present invention generally relates to systems and methods for preparing aliphatic alcohols. More particularly, embodiments described herein relate to systems and methods for preparing branched aliphatic alcohols using a hydrogenation unit and a dehydrogenation-isomerization unit.

Description of Related Art

Aliphatic alcohols are important compounds that may be used in a variety of applications or converted to other chemical compounds (e.g., surfactants, sulfates). Surfactants may be used in a variety of applications (e.g., detergents, soaps, oil recovery).

The structural composition of the aliphatic alcohol may influence the properties of the surfactant and/or detergent (e.g., water solubility, biodegradability and cold water detergency) produced from the aliphatic alcohol. For example, water solubility may be affected by the linearity of the aliphatic portion of the aliphatic alcohol. As the linearity of the aliphatic portion increases, the hydrophilicity (i.e., affinity for water) of the aliphatic alcohol surfactant may decrease. Thus, the water solubility and/or detergency performance of the aliphatic alcohol surfactant may decrease. Incorporating branches into the aliphatic portion of the aliphatic alcohol surfactant may increase the cold-water solubility and/or detergency of the aliphatic alcohol surfactant. Biodegradability, however, of the aliphatic alcohol surfactants may be reduced if the branches in the aliphatic portion of the alcohol surfactant include a high number of quaternary carbons. Incorporation of branches with a minimum number of quaternary carbon atoms into the aliphatic portion of the aliphatic alcohol surfactant may increase cold-water solubility and/or detergency of the alcohol surfactants while maintaining the biodegradability properties of the detergents.
The aliphatic portion of an aliphatic alcohol used to manufacture a surfactant may include one or more aliphatic alkyl groups as branches. Aliphatic alkyl groups that may form branches in the aliphatic portion may include methyl, ethyl, propyl or higher alkyl groups. Quaternary and tertiary carbons may be present when the aliphatic portion is branched. The number of quaternary and tertiary carbons may result from the branching pattern in the aliphatic portion. As used herein, the phrase “aliphatic quaternary carbon atom” refers to a carbon atom that is not bound to any hydrogen atoms.

Processes to manufacture branched primary alcohol compositions are described in U.S. Patent No. 5,849,960 to Singleton et al. entitled “Highly Branched Primary Alcohol Compositions, and Biodegradable Detergents Made Therefrom” and U.S. Patent No. 6,150,322 to Singleton et al., entitled “Highly Branched Primary Alcohol Compositions, and Biodegradable Detergents Made Therefrom.”

Summary of the Invention

In an embodiment, aliphatic alcohols may be produced by a method that includes hydrogenation of olefins to paraffins. A process feed stream entering a hydrogenation unit may include linear olefins and paraffins having an average carbon number from 7 to 18. In an embodiment, a process feed stream entering a hydrogenation unit includes linear olefins and paraffins having an average carbon number from 10 to 17. As used herein, the phrase “carbon number” refers to the total number of carbon atoms in a molecule. A process feed stream entering a hydrogenation unit is derived, in some embodiments from a Fischer-Tropsch process.

In the hydrogenation unit at least a portion of the olefins in the feed stream may be hydrogenated to form paraffins. The resulting paraffinic feed stream may be fed into a dehydrogenation-isomerization unit. At least a portion of the paraffins in the feed stream may be dehydrogenated to form olefins. The dehydrogenation-isomerization unit may also isomerize at least a portion of the resulting olefins and at least a portion of the olefins that were already present in the feed stream. The olefins produced from the dehydrogenation-isomerization unit may be hydroformylated to produce aliphatic alcohols. At least a portion of the aliphatic alcohols may have a branched aliphatic structure.

At least a portion of the paraffins in the feed stream may be dehydrogenated to form olefins in the dehydrogenation-isomerization unit. At least a portion of the resulting olefins and at least a portion of the olefins that were already present in the feed stream may also be
isomerized in the dehydrogenation-isomerization unit. An isomerization process converts linear olefins (e.g., unbranched olefins) into branched olefins. The isomerized olefins may be hydroformylated to produce aliphatic alcohols. After hydroformylation of the olefins, unreacted components from the hydroformylation process may be separated from the aliphatic alcohol products. Paraffins and unreacted olefins in the separated stream may be recycled back into the dehydrogenation-isomerization unit.

Process conditions in the dehydrogenation-isomerization unit may be such that the resulting branched olefins have an average number of branches per olefin molecule from about 0.7 to about 2.5. The branched olefins may include, but are not limited to, methyl and/or ethyl branched olefins. The isomerization process may produce branched olefins that include less than about 0.5 percent of quaternary aliphatic carbon atoms. The dehydrogenation-isomerization unit may include a catalyst that has two functions, to dehydrogenate the paraffins to olefins and to isomerize the olefins into branched olefins.

In an embodiment, a dehydrogenation-isomerization unit may include a plurality of zones. The plurality of zones may include a first reaction zone and a second reaction zone. The first reaction zone may be a dehydrogenation zone. The second reaction zone may be an isomerization zone. A hydrocarbon stream, containing olefins and paraffins, may enter the dehydrogenation zone. At least a portion of the paraffins in the hydrocarbon stream may be dehydrogenated to olefins to produce a stream enriched in olefins. The enriched olefin stream may be passed into the isomerization zone. In the isomerization zone, at least a portion of the olefins in the enriched olefin stream may be isomerized to branched olefins. The branched olefins may be converted to aliphatic alcohols by hydroformylation. After hydroformylation of the olefins, a paraffins and unreacted olefins stream may be separated from the produced aliphatic alcohol products. The paraffins and unreacted olefins stream may be recycled by directing at least a portion of the paraffins and unreacted olefins stream back into the dehydrogenation-isomerization unit and/or into a stream entering the dehydrogenation-isomerization unit.

In an embodiment, a dehydrogenation-isomerization unit may include a stacked bed configuration. The stacked bed may include a catalyst for dehydrogenation and a catalyst for isomerization of a hydrocarbon stream. At least a portion of the paraffins in the hydrocarbon stream may be dehydrogenated to olefins to produce a stream enriched in olefins. The enriched olefin stream may be passed into an isomerization zone. In the
isomerization zone, at least a portion of the olefins in the enriched olefin stream may be isomerized to branched olefins. The branched olefins may be converted to aliphatic alcohols by hydroformylation. After hydroformylation of the olefins, a paraffins and unreacted olefins stream may be separated from the produced aliphatic alcohol products.

The paraffins and unreacted olefins stream may be recycled by directing at least a portion of the paraffins and unreacted olefins stream back into the dehydrogenation-isomerization unit and/or into a stream entering the dehydrogenation-isomerization unit.

In certain embodiments, at least a portion of the aliphatic alcohols may be sulfated to form aliphatic sulfates. In some embodiments, aliphatic sulfates may include branched alkyl groups.

In certain embodiments, at least a portion of the produced aliphatic alcohols may be oxyalkylated to form oxyalkyl alcohols. In some embodiments, oxyalkyl alcohols may include branched alkyl groups. In some embodiments, at least a portion of the produced branched aliphatic alcohols may be ethoxylated to form branched ethoxyalkyl alcohols. At least a portion of the oxyalkyl alcohols may be sulfated to from oxyalkyl sulfates. In some embodiments, oxyalkyl sulfates may include branched alkyl groups.

**Brief Description of the Drawings**

Advantages of the present invention will become apparent to those skilled in the art with the benefit of the following detailed description of embodiments and upon reference to the accompanying drawings, in which:

FIG. 1 depicts a schematic diagram of an embodiment of a system for producing aliphatic alcohols using a hydrogenation unit and a dehydrogenation-isomerization unit;

FIG. 2 depicts a schematic diagram of an embodiment of a system for producing branched aliphatic alcohols using a hydrogenation unit, a dehydrogenation-isomerization unit and a separation unit to separate branched olefins from linear olefins and paraffins.

FIGS. 3 A-B depict schematic diagrams of embodiments of a system for producing branched aliphatic alcohols using a hydrogenation unit and a two-zone dehydrogenation-isomerization unit.

FIG. 3 depicts a schematic diagram of an embodiment of a system for producing branched aliphatic alcohols using a hydrogenation unit and a dehydrogenation-isomerization unit with a stacked bed catalyst configuration;
While the invention is susceptible to various modifications and alternative forms, specific embodiments thereof are shown by way of example in the drawing and will herein be described in detail. It should be understood that the drawings and detailed description thereto are not intended to limit the invention to the particular form disclosed, but on the contrary, the intention is to cover all modifications, equivalents, and alternatives falling within the spirit and scope of the present invention as defined by the appended claims.

**Detailed Description of the Embodiments**

Hydrocarbon products may be synthesized from synthesis gas (i.e., a mixture of hydrogen and carbon monoxide) using a Fischer-Tropsch process. Synthesis gas may be derived by partial combustion of petroleum (e.g., coal, hydrocarbons), by reforming of natural gas or by partial oxidation of natural gas. The Fischer-Tropsch process catalytically converts synthesis gas into a mixture of products that includes saturated hydrocarbons, unsaturated hydrocarbons and a minor amount of oxygen-containing products. The products from a Fischer-Tropsch process may be used for the production of fuels (e.g., gasoline, diesel oil), lubricating oils and waxes.

Fischer-Tropsch process streams may also be used to prepare commodity products, which have economic value. For example, linear olefins are commodity products that are useful for the production of surfactants. Using a portion of the process stream to produce linear olefins may increase the economic value of a Fischer-Tropsch process stream.

Surfactants derived from branched olefins may have different properties than surfactants derived from linear olefins. For example, surfactants derived from branched olefins may have increased water solubility and/or improved detergency properties compared to surfactants derived from linear olefins. Biodegradable properties of the surfactant, however, may be affected by the presence of quaternary carbon atoms in the branched portion of the surfactant. Surfactants made from branched olefins with a minimum number of quaternary carbon atoms may have similar biodegradable properties to surfactants derived from linear olefins. Production of branched olefins from a Fischer-Tropsch process stream may increase the economic value of the stream. In some embodiments, linear olefins may be converted into branched olefins with a minimum number of quaternary carbon atoms using an isomerization catalyst. Increasing the amount of branched olefins derived from a Fischer-Tropsch process stream may increase the economic value of the process streams.
Methods are described for increasing the amount of branched olefins derived from a process stream that includes certain amount of olefins, thus increasing the economic value of the process stream. Such methods are useful for both Fischer-Tropsch process streams and product streams from other sources that include hydrocarbons.

A hydrocarbon feed stream composition may include paraffins and olefins. At least a portion of the hydrocarbon stream may be made up of linear paraffins and olefins having at least 4 carbon atoms and up to 18 carbon atoms. A hydrocarbon feed stream may be obtained from a Fischer-Tropsch process or from an ethylene oligomerization process. Fischer-Tropsch catalysts and reaction conditions may be selected to provide a particular mix of products in the reaction product stream. For example, a Fischer-Tropsch catalyst and reaction conditions may be selected to increase the amount of olefins and decrease the amount of paraffins and oxygenates in the stream. Alternatively, the catalyst and reaction conditions may be selected to increase the amount of paraffins and decrease the amount of olefins and oxygenates in the stream.

The catalyst used in a Fischer-Tropsch process may be Mo, W, Group VIII compounds or combinations thereof. Group VIII compounds include, but are not limited to, iron, cobalt, ruthenium, rhodium, platinum, palladium, iridium and osmium. Combinations of Mo, W and Group VIII compounds may be prepared in the free metal form. In an embodiment, combinations of Mo, W and Group VIII compounds may be formed as alloys. Combinations of Mo, W and Group VIII compounds may be formed, in some embodiments, as oxides, carbides or other compounds. In other embodiments, combinations of Mo, W and Group VIII compounds may be formed as salts. Iron based and cobalt based catalysts have been used commercially as Fischer-Tropsch catalysts. Ruthenium catalysts tend to favor the formation of high melting waxy species under high-pressure conditions. Synthetic Fischer-Tropsch catalysts may include fused iron. In some embodiments, a fused iron Fischer-Tropsch catalyst may include a promoter (e.g., potassium or oxides on a silica support, alumina support or silica-alumina support). Cobalt metal may also be used in a Fischer-Tropsch catalyst. With the proper selection of supports, promoters and other metal combinations, a cobalt catalyst may be tuned to manufacture a composition enriched in the desired hydrocarbon species. Other catalysts, such as iron-cobalt alloy catalysts, are known for their selectivity toward the production of
olefins. Catalysts and combinations for manufacture of hydrocarbon species by a Fischer-Tropsch process are generally known.

While reference is made to a Fischer-Tropsch stream, any stream of olefins and saturated hydrocarbons may be suitable. Many Fischer-Tropsch streams may contain from 5 percent to 80 percent olefins, the remainder being saturated hydrocarbons comprising paraffins and other compounds.

In some embodiments, feed streams containing olefins and paraffins are obtained through cracking of paraffin wax or the oligomerization of olefins. Commercial olefin products manufactured by ethylene oligomerization are marketed in the United States by Chevron Phillips Chemical Company, Shell Chemical Company (as NEODENE®) and by British Petroleum. Cracking of paraffin wax to produce alpha-olefin and paraffin feed streams is described in U.S. Patent No. 4,579,986 to Sie, entitled “Process For The Preparation Of Hydrocarbons” and U.S. Patent Application Serial No. 10/153,955 of Ansorge et al., entitled “Process For The Preparation of linear Olefins and Use Thereof To Prepare Linear Alcohols.” Specific procedures for preparing linear olefins from ethylene are described in U.S. Patent No. 3,676,523 to Mason, entitled “Alpha-Olefin Production;” U.S. Patent No. 3,686,351 to Mason, entitled “Alpha-Olefin Production;” U.S. Patent No. 3,737,475 to Mason, entitled “Alpha-Olefin Production” and U.S. Patent No. 4,020,121 to Kister et al., entitled “Oligomerization Reaction System.” Most of the above-mentioned processes produce alpha-olefins. Higher linear internal olefins may be commercially produced (e.g., chlorination-dehydrochlorination of paraffins, paraffin dehydrogenation, isomerization of alpha-olefins).

In an embodiment, a feed stream is processed to produce a hydrocarbon stream that includes branched olefins. These branched olefins may be converted to branched aliphatic alcohols using various techniques. The feed stream may have a paraffin content range between about 50 percent by weight to about 90 percent by weight of the feed stream. In certain embodiments, a feed stream may have a paraffin content greater than about 90 percent by weight paraffins. The feed stream may also include olefins. The olefin content of the feed stream may be between about 10 percent by weight to about 50 percent by weight. In other embodiments, a feed stream may have an olefin content greater than 90 percent by weight olefins.
The composition of the feed stream may include hydrocarbons having an average carbon number ranging from 4 to 30. In an embodiment, an average carbon number of the hydrocarbons in a feed stream may range from 4 to 24. In other embodiments, an average carbon number of the feed stream may range from 4 to 18. An average carbon number of the hydrocarbons in a feed stream may range from 7 to 18. In certain embodiments, an average carbon number of the hydrocarbons in a feed stream may range from 10 to 17. In some embodiments, an average carbon number of hydrocarbons in a feed stream may range from 10 to 13. In other embodiments, an average carbon number of hydrocarbons in a feed stream may range from 14 to 17.

Branched chain olefins may be converted to branched aliphatic alcohols (e.g., branched primary alcohols) by a hydroformylation process. "Hydroformylation," as used herein, refers to the production of alcohols from olefins via a carbonylation and a hydrogenation process. Other processes may be used to produce aliphatic alcohols from olefins. Examples of other processes to produce aliphatic alcohols from olefins include, but are not limited to, hydration, oxidation and hydrolysis, sulfation and hydration, and epoxidation and hydration. The composition of an alcohol product stream may include aliphatic alcohols having an average carbon number ranging from 5 to 31. In an embodiment, an average carbon number of the aliphatic alcohols in an alcohol product stream may range from 7 to 20. In certain embodiments, an average carbon number of the aliphatic alcohols in an alcohol product stream may range from 11 to 18. In some embodiments, an average carbon number of aliphatic alcohols in an alcohol product stream may range from 11 to 14. In other embodiments, an average carbon number of aliphatic alcohols in an alcohol product stream may range from 15 to 18.

In certain embodiments, a first hydrocarbon stream may contain unwanted compounds (e.g., oxygenates and dienes) that may reduce catalyst selectivity in processes used to produce aliphatic alcohols. Removal of the unwanted compounds may be performed by hydrogenation of the first hydrocarbon stream. Hydrogenation of the first hydrocarbon stream, in certain embodiments, may produce a hydrocarbon stream that includes greater than about 90 percent paraffins. The hydrogenated hydrocarbon stream may be dehydrogenated to produce an olefinic stream. The catalyst used in the dehydrogenation process may control the position of the olefin double bond. In certain embodiments, an olefinic hydrocarbon stream may include olefins in which greater than 70
percent of the olefins are alpha-olefins of a linear carbon skeletal structure. In other embodiments, an olefinic hydrocarbon stream may include olefins in which 50 percent or more of the olefin molecules present may be internal olefins.

A first hydrocarbon stream may be introduced into hydrogenation unit 110 via first conduit 112 as depicted for System 100, in FIG. 1. The first hydrocarbon stream includes olefins and paraffins. In hydrogenation unit 110, at least a portion of the olefins in the first hydrocarbon stream may be hydrogenated to paraffins to produce a second hydrocarbon stream.

Reaction conditions in hydrogenation unit 110 may be controlled to hydrogenate olefins and dienes and to remove oxygenates. An operating temperature of hydrogenation unit 110 may range between about 100 °C and about 300 °C. In some embodiments, an operating temperature may range from about 150°C to about 275 °C. In other embodiments, an operating temperature may range from about 175 °C to 250 °C. An operating pressure may range from about 5 atmospheres (506 kPa) to about 150 atmospheres (1520 kPa). In some embodiments, an operating pressure may range from 10 atmospheres psi (1013 kPa) to about 50 atmospheres (5065 kPa).

Hydrogenation processes may be carried out using any type of catalyst bed arrangement (e.g., fluidized bed, moving bed, slurry phase bed or a fixed bed). In certain embodiments, a fixed bed arrangement may be used. In a fixed bed system, hydrogen may be supplied to the hydrogenation stage at a gas hourly space velocity in the range from about 100 normal liter gas/liter catalyst/hour (NL/L/hr) to about 1000 NL/L/hr. In some embodiments, hydrogen may be supplied at a gas hourly space velocity in the range from about 250 NL/L/hr to 5000 NL/L/hr. “Gas space velocity as expressed in units of normal liter of gas/liter of catalyst/hour,” as used herein, is the volume of a gas in liters at standard conditions of 0 °C and 760 mm Hg.

Hydrogenation catalysts are generally known and are commercially available in a large variety of compositions. In some embodiments, a hydrogenation catalyst may include one or more metals from Groups VIB and VII of the periodic Table of the Elements. In certain embodiments, metals may include, but are not limited to, molybdenum, tungsten, cobalt, nickel, ruthenium, iridium, osmium, platinum and palladium. The hydrogenation catalyst may include a refractory oxide or a silicate as a binder.
Hydrogenation reaction conditions and catalysts are described in European Patent No. 0 583 836 to Eilers et al., entitled “Process For The Preparation of Hydrocarbon Fuels;” European Patent No. 0 668 342 to Eilers et al., entitled “Lubricating Base Oil Preparation Process.” Hydrogenation reaction conditions and catalysts are also described in U.S. Patent No. 5,371,308 to Gosselink et al., entitled “Process For The Preparation Of Lower Olefins.”

In certain embodiments, a second hydrocarbon stream that includes paraffins and olefins may be introduced into a dehydrogenation-isomerization unit. The dehydrogenation-isomerization unit may replace two independent units (e.g., an isomerization unit and a dehydrogenation unit). The dehydrogenation-isomerization unit may dehydrogenate paraffins to olefins and isomerize the resulting olefins and/or initial olefins present in the hydrocarbon stream to branched olefins. In an embodiment, a catalyst may perform the dehydrogenation-isomerization of the hydrocarbons in the second hydrocarbon stream. In certain embodiments, a catalyst may be a single catalyst. The catalyst, in some embodiments, may be a mixture of two catalysts (e.g., a dehydrogenation catalyst and an isomerization catalyst). In other embodiments, two separate catalysts located in different zones or in a stacked bed configuration in one dehydrogenation-isomerization unit may perform the dehydrogenation-isomerization process. As used herein, “a dehydrogenation-isomerization catalyst” may be one or more catalysts.

In certain embodiments, a dehydrogenation-isomerization unit may have several points of entry to accommodate different process streams. The process streams may be from other processing units and/or storage units. Examples of process streams include, but are not limited to, a diluent hydrocarbon stream, and/or other hydrocarbon streams that include olefins and paraffins derived from other processes. As used herein, “entry into the dehydrogenation-isomerization unit” refers to entry of process streams into the dehydrogenation-isomerization unit through one or more entry points.

A second hydrocarbon stream, including a mixture of olefins and paraffins, may be introduced into dehydrogenation-isomerization unit 114 via second conduit 116. In dehydrogenation-isomerization unit 114, at least a portion of the paraffins in the second hydrocarbon stream may be dehydrogenated to olefins. At least a portion of the resulting olefins and at least a portion of the olefins that were already present in the feed stream may
be isomerized to produce a third hydrocarbon stream. The isomerization process converts linear olefins (i.e., unbranched olefins) into branched olefins.

The catalyst used for the dehydrogenation-isomerization of the second hydrocarbon stream may be based on a zeolite catalyst modified with one or more metals or metal compounds. The catalyst used in dehydrogenation-isomerization unit 114 to treat the olefins in the second hydrocarbon stream may be effective for skeletally isomerizing linear olefins in the process stream into olefins having an average number of branches per olefin molecule chain greater than about 0.7. In certain embodiments, an average number of branches per olefin molecule chain may range from about 0.7 to about 2.5. In some embodiments, an average number of branches per olefin molecule chain may range from about 0.7 to about 2.2. In other embodiments, an average number of branches per olefin molecule chain may range from about 1.0 to about 2.2.

The dehydrogenation-isomerization catalyst may contain a zeolite having at least one channel with a crystallographic free channel diameter greater than about 4.2 Å and less than about 7 Å, measured at room temperature. As used herein, “channel diameter or size” refers to an effective channel diameter or size for diffusion. The zeolite may have no channels present that have a free channel diameter greater than about 7 Å. The catalyst may contain at least one channel having a crystallographic free diameter at the entrance of the channel greater than about 4.2 Å and less than about 7 Å. The catalyst may not have a channel with a diameter at the entrance, which exceeds the 7 Å upper limit of the range. Zeolites possessing channel diameters greater than about 7 Å may be susceptible to undesirable olefin by-products (e.g., aromatization, oligomerization, alkylation, coking). In some embodiments, a zeolite may not contain a channel having a free diameter along either of the x or y planes of greater than about 4.2 Å. A small channel size may prevent diffusion of the olefin into and/or out of the channel pore once the olefin becomes branched. A zeolite may have at least one channel with a free diameter of the channel within a range of greater than about 4.2 Å and less than about 7 Å.

In an embodiment, an olefin molecule, due to its high carbon chain length, may not have to enter into the zeolite channel, diffuse through, and exit the other end of the channel. The rate of branching seen when passing the olefin across the zeolite may not correspond to the theoretical rate of branching if each olefin molecule were to pass through the channels. Most of the olefins may partially penetrate the channel for a distance effective to branch the
portion of the chain within the channel and subsequently withdraw from the channel once isomerized. In an embodiment of a method to produce aliphatic alcohols, olefin molecules in a hydrocarbon stream may predominately have a structure which is branched at the ends of the olefin carbon backbone, and substantially linear towards the center of the molecule, (e.g., at least 25 percent of the carbons at the center are unbranched).

In certain embodiments, a zeolite catalyst structure may contain channels having free diameters greater than about 4.2 Å and less than about 7 Å along both the x and y planes in the [001] view. Zeolites with the specified channel size may be referred to as medium or intermediate channel zeolites and typically have a 10-T member (or puckered 12-T member) ring channel structure in one view and a 9-T member or less (small pore) in another view, if any. There is no limit to channel numbers or orientation (e.g., parallel, non-interconnecting intersections, or interconnecting at any angle) in the zeolite.

Examples of zeolites with a channel size from about 4.2 Å to 7.0 Å include molecular sieves, ferrierite, A1PO-31, SAPO-11, SAPO-31, SAPO-41, FU-9, NU-10, NU-23, ZSM-12, ZSM-22, ZSM-23, ZSM-35, ZSM-48, ZSM-50, ZSM-57, SUZ-4A, MeAPO-11, MeAPO-31, MeAPO-41, MeAPSO-11, MeAPSO-31, and MeAPSO-41, MeAPSO-46, ELAPO-11, ELAPO-31, ELAPO-41, ELAPSO-11, ELAPSO-31, and ELAPSO-41, laumontite, cancrinite, offretite, hydrogen form of stilbite, the magnesium or calcium form of mordenite and partheite. The isotypic structures of the zeolite frameworks, known under other names, may be considered equivalent. Zeolite framework is described by Flanigen et al.; in "Aluminophosphate Molecular Sieves and the Periodic Table," New Developments in Zeolite Science Technology, 1986, Kodansha Ltd., Tokyo, Japan.

Many natural zeolites such as ferrierite, heulandite and stilbite may feature a one-dimensional pore structure with a pore size at or slightly smaller than about 4.2 Å in diameter. U.S. Patent No. 4,795,623 to Evans, entitled "Time Effective Method For Preparing Ferrierite" and U.S. Patent No. 4,942,027 to Evans, entitled "Method for Preparing Ferrierite," describe converting channels in natural zeolites to larger channels. Channels in natural zeolites may be converted to zeolites with desired larger channel sizes by removing an associated alkali metal or alkaline earth metal by generally known methods (e.g., ammonium ion exchange, optionally followed by calcination, to yield a zeolite in substantially a hydrogen form). Replacing the associated alkali or alkaline earth metal with the hydrogen form may enlarge the channel diameter. In some embodiments, natural
zeolites (e.g., some forms of mordenite) may have a channel size greater than 7 Å. The channel size may be reduced by substituting an alkali metal for larger ions (e.g., a larger alkaline earth metal).

In certain embodiments, zeolites may have a ferrierite isotypic (or homeotypic) framework structure. The prominent structural features of ferrierite found by x-ray crystallography may be parallel channels in the alumino-silicate framework. The parallel channels may have an elliptical cross section. Zeolites having a ferrierite isotypic framework structure are described in European Patent No. 55 529 to Seddon et al., entitled, “Zeolites,” and European Patent No. 103 981 to Whittam, entitled “Zeolites.” Zeolites having a ferrierite isotypic framework are also described in U.S. Patent No. 4,016,245 to Plank et al., U.S. Patent No. 4,578,259 to Morimoto et al., entitled “Process For Preparing A Crystalline Aluminosilicate;” entitled “Crystalline Zeolite And Method Of Preparing Same” and U.S. Patent No. 4,375,573 to Young et al., entitled “Selective Production And Reaction of P-Disubstituted Aromatics Over Zeolite ZSM-48.”

In an embodiment, a hydrogen form of ferrierite (H-ferrierite) may be considered to be substantially one-dimensional. H-ferrierite may have parallel running channels. H-ferrierite may have elliptical channels that have free diameters of 4.2 Å by 5.4 Å along the x and y planes in the [001] view. The channels may be large enough to permit entry of a linear olefin and diffusion out of or through the channel of the methyl branched isoolesfin. The channels may be small enough to retard coke formation. Methods for preparing various H-ferrierite are described in U.S. Patent No. 5,985,238 to Pasquale et al., entitled “Process For Preparing Ferrierite;” U.S. Patent No. 4,251,499 to Namne et al., entitled “Process For The Preparation Of Ferrierite;” U.S. Patent No. 4,795,623 to Evans, entitled “Time Effective Method For Preparing Ferrierite” and U.S. Patent No. 4,942,027 to Evans, entitled “Method for Preparing Ferrierite.”

In certain embodiments, a dehydrogenation-isomerization catalyst may be combined with a refractory oxide that serves as a binder material. Suitable refractory oxides include, but are not limited to, natural clays (e.g., bentonite, montmorillonite, attapulgite, and kaolin), alumina, silica, silica-alumina, hydrated alumina, titania, zirconia or mixtures thereof.

Examples of alumina binders may include, but are not limited to, pseudoboehmite, gamma and bayerite aluminas. Alumina binders may be commercially available (e.g.,
LaRoche Chemicals manufactures VERSAL® aluminas and Sasol manufactures CATAPAL® aluminas. In an embodiment, high-dispersity alumina powders may be used as alumina binders when extrusion is utilized for catalyst preparation. High-dispersity alumina powders may have a dispersity of greater than 50 percent in an aqueous acid dispersion having an acid content of 0.4-milligram equivalents of acid (acetic) per gram of powder. Such high-dispersity aluminas may be exemplified by CATAPAL® alumina manufactured by Sasol.

A weight ratio of zeolite to binder material may range from about 10:90 to about 99.5:0.5. In some embodiments, a weight ratio may range from about 75:25 to about 99:1. In other embodiments, a weight ratio of zeolite to binder material may range from about 80:20 to about 98:2. In certain embodiments, a weight ratio of zeolite to binder material may range from about 85:15 to about 95:5 on an anhydrous basis.

In certain embodiments, a dehydrogenation-isomerization catalyst may be prepared with one or more monocarboxylic acids and/or inorganic acids. In addition to the monocarboxylic and/or inorganic acids, at least one organic acid with at least two carboxylic acid groups ("polycarboxylic acid") may be used. Monocarboxylic acids may have a substituted or unsubstituted hydrocarbyl group having 1 to 20 carbon atoms. The hydrocarbyl group may be aliphatic, cyclic or aromatic. Examples of monocarboxylic acids having 1 to 20 carbon atoms include, but are not limited to, acetic acid, formic acid, propionic acid, butyric acid, caproic acid, glycolic acid, lactic acid, hydroxybutyric acid, hydroxycyclopentanoic acid, salicylic acid, mandelic acid, benzoic acid and fatty acids. Examples of inorganic acids include, but are not limited to, nitric acid, phosphoric acid, sulfuric acid and hydrochloric acid.

The polycarboxylic acid may, in certain embodiments, be an organic acid with two or more carboxylic acid groups attached through a carbon-carbon bond linkage to a hydrocarbon segment. The linkage may be at any portion of the hydrocarbon segment. The polycarboxylic acid may have a hydrocarbon segment with less than 10 carbon atoms. The hydrocarbon segment may be aliphatic, cyclic or aromatic. The hydrocarbon segment may have zero carbon atoms for oxalic acid with two carboxylic acid groups attached through the carbon-carbon bond. Examples of the polycarboxylic acids include, but are not limited to, tartaric acid, citric acid, malic acid, oxalic acid, adipic acid, malonic acid, galactaric acid, 1,2-cyclopentane dicarboxylic acid, maleic acid, fumaric acid, itaconic acid, phthalic acid.
acid, terephthalic acid, phenylmalonic acid, hydroxyphthalic acid, dihydroxyfumaric acid, tricarballylic acid, benzene-1,3,5-tricarboxylic acid, isocitric acid, mucic acid and glucaric acid. The polycarboxylic acids may be any isomers of the above acids. In some embodiments, the polycarboxylic acids may be any stereoisomers of the above acids. In an embodiment, polycarboxylic acids with at least two carboxylic acid groups and at least one hydroxyl group are used. In an embodiment, citric acid, tartaric acid and malic acid may be used as polycarboxylic acids.

Metals incorporated into a dehydrogenation-isomerization catalyst may be metals that promote the oxidation of coke in the presence of oxygen at a temperature greater than about 250 °C and the dehydrogenation of paraffins. "Metal(s)," as used herein, refers to metals of a zero oxidation state and/or higher oxidation states (e.g., metal oxides). As used herein, "coke" refers to a product from thermal degradation of larger molecules into smaller molecules.

Metals used in the dehydrogenation-isomerization catalyst may be transition and rare earth metals. Coke oxidation-promoting metals include, but are not limited to, Groups IB, VB, VIB, VIIB, VIII of the transition metal series of the Periodic Table and/or combinations thereof. In certain embodiments, Pd, Pt, Ni, Co, Mn, Ag, Cr and/or combinations thereof may be used in the dehydrogenation-isomerization catalyst. In other embodiments, metals oxides such as, but not limited to, chrome oxide, iron oxide, noble metals, or mixtures thereof may be used as coke-oxidizing compounds in the catalyst.

An amount of the metal introduced may range from about 5 parts per million ("ppm") up to about 15 percent by weight. In certain embodiments, an amount of metal may range from about 5 ppm to about 10 percent by weight. In some embodiments, an amount of metal may range from about 5 ppm to about 5 percent by weight.

Noble metals (e.g., platinum and/or palladium) may be used in smaller amounts of metals than other metals incorporated into a zeolite and/or binder. "Noble metals," as used herein, refers to metals of the group that includes platinum, palladium, iridium, ruthenium, osmium and rhodium. In certain embodiments, an amount of noble metal(s) may range from about 5 ppm to about 2 percent by weight, basis metal, of the final catalyst. In some embodiments, an amount of noble metal(s) may range from about 5 ppm to about 1000 ppm, basis metal, of the final catalyst. In other embodiments, an amount of noble metal(s) may range from about 5 ppm to about 3000 ppm, basis metal, of the final catalyst. An
amount of noble metal(s) used in a dehydrogenation-isomerization catalyst may, in certain embodiments, range from about 5 ppm to about 2000 ppm by weight, basis metal, of the final catalyst. An amount of noble metal(s) sufficient to promote regeneration without deteriorating the performance of the catalyst may be in the range from about 30 ppm to about 100 ppm. Higher amounts of platinum and/or palladium (e.g., greater than about 2% by weight) may have an adverse effect on the catalyst (e.g., run life, olefin isomerization activity, selectivity).

In an embodiment, zeolite powder and alumina powder may be mixed (e.g., mulled) with water and one or more metal compounds of the catalyst. The resulting mixture may be formed into a pellet. Catalysts prepared by mulling may have superior olefin isomerization performance over catalysts prepared by impregnation. The term "mulling," as used herein, refers to mixing of powders to which sufficient water has been added to form a thick paste and wherein the mixing is accompanied by concomitant shearing of the paste. Commercially available mullers such as the Lancaster Mix Muller and the Simpson Mix Muller may be used.

The pellet may be formed, in some embodiments, by extrusion. One or more peptizing acid (e.g., nitric acid, acetic acid, citric acid or mixtures thereof) may be added to the mixture and optional extrusion aids such as cellulose derivatives (e.g., METHOCEL®P4M, hydroxypropyl methylcellulose, manufactured by The Dow Chemical Company) may be utilized. The amounts of peptizing acid used may be determined by routine experimentation to provide a plastic, extrudable material. The term "pellets," as used herein, refers to any shape or form of consolidated materials.

In certain embodiments, a noble metal such as platinum and/or palladium may be added to the zeolitic catalyst after pelletization. Common metal incorporation methods known to those skilled in the art (e.g., impregnation, noble metal ion exchange and co-mulling) may be used to produce a working catalyst useful in dehydrogenation-isomerization of paraffins. The addition of noble metals to the catalyst may aid in the dehydrogenation reaction of paraffins. Pellets containing noble metals may be calcined at a temperature range from about 250 °C to about 700 °C. In certain embodiments, a calcination temperature may range from about 300°C to about 600 °C. In some embodiments, a calcination temperature may range from about 450 °C to about 525 °C.
The dehydrogenation-isomerization catalyst may be contacted with the second hydrocarbon stream in dehydrogenation-isomerization unit 114 under a variety of conditions to dehydrogenate at least a portion of the paraffins to olefins and isomerize at least a portion of the resulting olefins. In dehydrogenation-isomerization unit 114, reaction temperatures may range from about 300 °C to about 700 °C. A reaction temperature, in some embodiments, may range from about 350 °C to about 550 °C. A total pressure of dehydrogenation-isomerization unit 114 during the reaction may range from about 0.010 atmosphere (1 kPa) to about 25.0 atmospheres (2534 kPa). In an embodiment, a total pressure of dehydrogenation-isomerization unit 114 during the reaction may range from about 0.010 atmosphere (1 kPa) to about 15.0 atmospheres (1520 kPa). In other embodiments, a total pressure of dehydrogenation-isomerization unit 114 during the reaction may range from about 1 atmosphere (101 kPa) to about 5.0 atmospheres (507 kPa). In order to prevent coking, hydrogen may be fed together with the second hydrocarbon stream. Hydrogen gas and paraffins present in the second hydrocarbon stream may be fed at a hydrogen to paraffin molar ratio in the range from about 0.1 to about 20. In certain embodiments, a hydrogen to paraffin molar ratio may be in the range from about 1 to about 10.

Residence time in dehydrogenation-isomerization unit 114 may be selected such that conversion level of the paraffins to olefins may be kept below 40 mole percent. In an embodiment, a conversion level ranges from 5 mole percent to 30 mole percent. By keeping the conversion level low, side reactions (e.g., diene formation and cyclization reactions) may be minimized. Olefin conversion may be increased by varying the reaction conditions (e.g., temperature, residence time) as long as side reactions remain below acceptable limits. Olefins produced in dehydrogenation-isomerization unit 114 may have a higher degree of branching than a paraffin feed to the dehydrogenation-isomerization unit. It should be understood that the concentration of olefins produced via dehydrogenation-isomerization unit 114 may be limited by the thermodynamic equilibrium of olefins and paraffins at the reaction temperature. The conditions for olefin isomerization dehydrogenation-isomerization 114 may be controlled such that the number of carbon atoms in the olefins prior to and subsequent to the isomerization conditions is substantially the same.
Branched olefins produced in dehydrogenation-isomerization unit 114 may include methyl, ethyl and/or longer carbon chain branches. Hydrogen Nuclear Magnetic Resonance (\( ^1H \) NMR) analysis of the isomerized olefin composition may be performed. Branched olefins may include quaternary and/or tertiary aliphatic carbons. In certain embodiments, an amount of quaternary aliphatic carbons produced in a dehydrogenation-isomerization unit may be minimized. \( ^1H \) NMR analysis of the olefins may indicate the extent of isomerization of the olefins in the hydrocarbon stream. \( ^1H \) NMR analysis may be capable of differentiating a wide range of olefin structures.

In an embodiment, an average number of branches per olefin molecule present in the produced branched olefin composition may be greater than 0.7. In certain embodiments, an average number of branches per olefin molecule present in the branched olefin composition is from about 0.7 to about 2.5. In some embodiments, an average number of branches per olefin molecule present in the branched olefin composition is from about 0.7 to about 2.2. In certain embodiments, an average number of branches per olefin molecule present in the branched olefin composition is from about 1.0 to about 2.2. The degree of branching in the product may be controlled by controlling process conditions used in a dehydrogenation-isomerization unit. For example, high reaction temperatures and lower feed rates may result in a higher degree of branching. Methyl branches may represent between about 20 percent to about 99 percent of the total number of branches present in the olefin molecules. In some embodiments, methyl branches may represent greater than about 50 percent of the total number of branches in the olefin molecules. The number of ethyl branches in the olefin molecules may represent, in certain embodiments, less than about 30 percent of the total number of branches. In other embodiments, a number of ethyl branches, if present, may be between about 0.1 percent and about 2 percent of the total number of branches. Branches other than methyl or ethyl, if present, may be less than about 10 percent of the total number of branches.

Aliphatic quaternary carbon atoms present in the branched olefin composition may be less than about 2 percent of the carbon atoms present. In an embodiment, a number of aliphatic quaternary carbon atoms present is less than about 1 percent of the carbon atoms present. For applications in which biodegradability is important, the number of aliphatic quaternary carbon atoms may be less than about 0.5 percent of the carbon atoms present. In an embodiment, a number of aliphatic quaternary carbon atoms is less than about 0.3
percent of the carbon atoms present. In other embodiments, a number of aliphatic quaternary carbon atoms present in the branched olefin composition is between about 0.01 percent and about 0.3 percent of the aliphatic carbon atoms present.

A third hydrocarbon stream may exit dehydrogenation-isomerization unit 114 and be transferred to other processing units (e.g., a hydroformylation unit, separation units, an alkylation units) via third conduit 118. At least a portion of the third hydrocarbon stream may exit dehydrogenation-isomerization unit 114 and be introduced into hydroformylation unit 120 via third conduit 118. In hydroformylation unit 120, at least a portion of the olefins in the third hydrocarbon stream may be converted to alcohols. At least a portion of the produced alcohols and at least a portion of the unreacted components of the third hydrocarbon stream may form a hydroformylation reaction stream.

In an embodiment, olefins may be separated, if desired, from the third hydrocarbon stream through techniques generally known in the art (e.g., distillation, molecular sieves, extraction, adsorption, adsorption/desorption, and/or membranes). Separation of at least a portion of the branched olefins from the linear olefins and paraffins may increase the concentration of branched olefins entering the hydroformylation unit. In addition, separation of at least a portion of the branched olefins from the linear olefins and paraffins may influence the ratio of linear to branched olefins produced in the hydroformylation unit.

Referring to FIG. 2, a third hydrocarbon stream may exit dehydrogenation-isomerization unit 114 and enter separation unit 122 via separation conduit 124. Separation unit 122 may produce at least two streams, a branched olefins stream and a linear olefins and paraffins stream. In separation unit 122, the third hydrocarbon stream may be contacted with organic and/or inorganic molecular sieves (e.g., zeolite or urea) with the correct pore size for branched olefins and/or linear olefins and paraffins. Subsequent desorption (e.g., solvent desorption) of at least a portion of the branched olefins and/or at least a portion of the linear olefins and paraffins from the molecular sieves may produce at least two streams (e.g., a branched olefins stream and a linear olefins and paraffins stream).

Separation unit 122 may include a fixed bed containing adsorbent for separation of the third hydrocarbon stream to produce a branched olefin and paraffins stream and a linear olefins and paraffins stream. Separation temperatures in separation unit 122 may range from about 100 °C to about 400 °C. In some embodiments, separation temperatures may range from 180 °C to about 380 °C. Separation in separation unit 122 may be conducted at
a pressure ranging from about 2 atmospheres (202 kPa) to about 7 atmospheres (710 kPa). In some embodiments, a pretreatment of a third hydrocarbon stream may be performed to prevent adsorbent poisoning. An example of an adsorption/desorption process is a Molex process using Sorbex® separations technology (UOP process, UOP, Des Plaines, IL).

Adsorption/desorption processes are described in U.S. Patent No. 6,225,518 to Sohn et al., entitled “Olefinic Hydrocarbon Separation Process;” U.S. Patent No. 5,292,990 to Kantner et al., entitled, “Zeolite Compositions For Use in Olefinic Separations” and U.S. Patent No. 5,276,246 to McCulloch et al., entitled “Process For Separating Normal Olefins From Non-Normal Olefins.”

At least a portion of the linear olefins and paraffins stream may be transported to other processing units and/or stored on site. In an embodiment, at least a portion of the linear olefins and paraffins stream may be combined with the first hydrocarbon stream in first conduit 112 via linear olefin and paraffin recycle conduit 126. The combined stream may enter hydrogenation unit 110 via first conduit 112 to continue the process to produce aliphatic alcohols. In some embodiments, a linear olefins and paraffins stream may be introduced directly into hydrogenation unit 110.

At least a portion of the branched olefins stream may be transported and utilized in other processing streams and/or stored on site via branched olefins conduit 128. In some embodiments, at least a portion of a branched olefins stream may exit separation unit 122 and be introduced into third conduit 118 via branched olefins conduit 128. In other embodiments, at least a portion of a branched olefins stream may exit separation unit 122 and be introduced directly into a hydroformylation unit.

Referring to FIG. 1, the third hydrocarbon stream may exit dehydrogenation-isomerization unit and enter hydroformylation unit 120 via third conduit 118.

Hydroformylation unit 120 may have several points of entry to accommodate entry of additional process streams. As used herein, “stream entering into the hydroformylation unit” is defined as the entry of process streams into the hydroformylation unit through one or more entry points. Examples of such process streams include, but are not limited to, additional streams from dehydrogenation-isomerization unit 114, a diluent hydrocarbon stream, gases and/or other hydrocarbon streams that include olefins and paraffins derived from other processes.
In a hydroformylation process, olefins are converted to aldehydes, alcohols or a combination thereof by reaction of at least a portion of the olefins with carbon monoxide and hydrogen according to an Oxo process. As used herein, an “Oxo process” refers to the reaction of an olefin with carbon monoxide and hydrogen in the presence of a metal catalyst (e.g., a cobalt catalyst) to produce an alcohol containing one more carbon atom than the starting olefin. In other hydroformylation processes, a “modified Oxo process” is used. As used herein, a “modified Oxo process” refers to an Oxo process that uses a phosphine, phosphite, arsine or pyridine ligand modified cobalt or rhodium catalyst.


A hydroformylation catalyst used in hydroformylation unit 130 may include a metal from Group VIII of the Periodic Table. Examples of Groups VIII metals include cobalt, rhodium, nickel, palladium or platinum. The Group VIII metal may be used as a complex compound. A complex compound may be a Group VIII metal combined with a ligand. Examples of ligands include, but are not limited to, a phosphine, phosphite, arsine, stibine or pyridine ligand. Examples of hydroformylation catalysts include, but are not limited to, cobalt hydrocarbonyl catalyst, cobalt-phosphine ligand catalyst, rhodium-phosphine ligand catalyst or combinations thereof.

In hydroformylation unit 120, olefins in the third hydrocarbon stream may be hydroformylated using a continuous, semi-continuous or batch process. In case of a continuous mode of operation, the liquid hourly space velocities may be in the range of about 0.1 h⁻¹ to about 10 h⁻¹. When operating hydroformylation unit 120 as a batch process, reaction times may vary from about 0.1 hours to about 10 hours or even longer. Reaction temperatures in hydroformylation unit 120 may range from about 100 °C to about 300 °C. In certain embodiments, reaction temperatures in the hydroformylation unit ranging from about 125 °C to about 250 °C may be used. Pressure in hydroformylation unit 120 may range from about 1 atmosphere (101 kPa) to about 300 atmospheres (30398 kPa). In an embodiment, a pressure from about 20 (2027 kPa) to about 150 atmospheres (15199 kPa) may be used. An amount of catalyst relative to the amount of olefin to be hydroformylated may vary. Typical molar ratios of catalyst to olefin in the third hydrocarbon stream may range from about 1:1000 to about 10:1. A ratio of between about 1:10 and about 5:1 may be used in certain embodiments. In an embodiment, a diluent stream may be added to hydroformylation unit 120 to control reaction conditions. The diluent stream may include solvents that do not interfere substantially with the desired reaction. Examples of such solvents include, but are not limited to, alcohols, ethers, acetonitrile, sulfolane and paraffins.

Mono-alcohol selectivities of at least 90 percent and even of at least 92 percent may be achieved in hydroformylation unit 120. In addition, olefin conversions to aliphatic alcohols may range from about 50 percent by weight to greater than about 95 percent by weight. In certain embodiments, olefin conversion to aliphatic alcohols may be greater
than 75 percent by weight. In some embodiments, olefin conversion to aliphatic alcohols may be greater than about 99 percent by weight.

Isolation of aliphatic alcohols produced from the hydroformylation reaction product stream may be achieved by generally known methods. In an embodiment, isolation of the aliphatic alcohols includes subjecting the produced aliphatic alcohols to a first distillation, a saponification, a water washing treatment and a second distillation.

The hydroformylation reaction mixture stream may enter separator 130 via fourth conduit 132. In separator 130, the hydroformylation reaction product stream may be subjected to a first distillation step (e.g., flash distillation or a short path distillation). In an embodiment, a short path distillation may be used to produce at least two streams, a bottom stream and a top stream. At least a portion of the bottom stream may be recycled to hydroformylation unit 120 via bottom stream recycle conduit 134, in certain embodiments. The top stream may include, but is not limited to, paraffins, unreacted olefins and a crude aliphatic alcohol product.

In an embodiment, a top stream may be subjected to a saponification treatment to remove any acids and esters present in the stream. Saponification may be performed by contacting the top stream with an aqueous solution of a hydroxide base (e.g., sodium hydroxide or potassium hydroxide) at elevated temperatures with agitation. The saponification may be carried out by contacting the top stream with an aqueous 0.5 percent to 10 percent hydroxide base solution at a crude alcohol/water ratio of 10:1 to 1:1. The amount of hydroxide base used may depend on an estimated amount of esters and acids present.

Saponification of the top stream may be carried out batch-wise or continuously. The top stream may be subjected to one or more saponification processes. Saponification reaction temperatures may be from about 40 °C to about 99 °C. In an embodiment, saponification temperatures may range from about 60 °C to about 95 °C. Mixing of the top stream with the basic water layer may be performed during the saponification reaction. Separation of the top stream from the basic water layer may be performed using known methods. The top stream may be subjected to a water wash after separation to remove any sodium salts present. The top stream may be separated using generally known techniques (e.g., fractional distillation) to produce at least two streams, a crude alcohol product stream and a paraffins and unreacted olefins stream. As used herein, “fractional distillation” refers
to the distillation of liquids and subsequent collection of fractions of liquids determined by boiling point. The paraffins and unreacted olefins stream may be recycled, transported to other units for processing, stored on site, transported offsite and/or sold.

In certain embodiments, a crude aliphatic alcohol product stream may contain unwanted by-products (e.g., aldehydes, hemi-acetals). The by-products may be removed by subjecting the crude alcohol product stream to a hydrofinishing treatment step to produce an aliphatic alcohol product stream. "Hydrofinishing," as used herein, refers to a hydrogenation reaction carried out under relatively mild conditions. Hydrofinishing may be carried out using conventional hydrogenation processes. Conventional hydrogenation processes may include passing the crude alcohol feed together with a flow of hydrogen over a bed of a suitable hydrogenation catalyst. The aliphatic alcohol product stream may include greater than about 50 percent by weight of the produced aliphatic alcohols. In some embodiments, the aliphatic alcohol product stream may include greater than 80 percent by weight of the produced aliphatic alcohols. In other embodiments, the aliphatic alcohol product stream may include greater than 95 percent by weight of the produced aliphatic alcohols. The aliphatic alcohol product stream may include branched aliphatic primary alcohols. The resulting aliphatic alcohols in the aliphatic alcohol product stream may be sold commercially, transported off-site, stored on site and/or used in other processing units via product conduit 136.

The composition of an aliphatic alcohol product stream may include hydrocarbons with an average carbon number ranging from 8 to 19. In an embodiment, an average carbon number of the hydrocarbons in aliphatic alcohol product stream may range from 10 to 17. In certain embodiments, an average carbon number of the feed stream may range from 10 to 13. In other embodiments, an average carbon number of the feed stream may range from 14 to 17. In some embodiments, branched primary alcohol products may be used as the precursor for the manufacture of anionic sulfates, including aliphatic sulfates and oxyalkyl sulfates and oxyalkyl alcohols.

Aliphatic alcohols may have slightly higher aliphatic branching and slightly higher number of quaternary carbons as the olefin precursor. In some embodiments, aliphatic branching may include methyl and/or ethyl branches. In other embodiments, aliphatic branching may include methyl, ethyl and higher aliphatic branching. In certain embodiments, a number of quaternary carbon atoms in the aliphatic alcohol product may be
less than 0.5. In other embodiments, a number of quaternary carbon atoms in the aliphatic alcohol product may be less than 0.3. Branching of the alcohol product may be determined by $^1$H NMR analysis. The number of quaternary carbon atoms may be determined by $^{13}$C NMR. A $^{13}$C NMR method for determining quaternary carbon atoms for branched aliphatic alcohols is described in U.S. Patent No. 6,150,322 to Singleton et al., entitled, “Highly Branched Primary Alcohol Compositions and Biodegradable Detergents Made Therefrom.”

In certain embodiments, at least a portion of the paraffins and unreacted olefins stream may be combined with the second hydrocarbon stream in second conduit 116 to produce a combined stream via fifth conduit 138. The combined stream may be introduced into dehydrogenation-isomerization unit 114 via second conduit 116. At least a portion of the olefins in the combined stream may be isomerized to branched olefins. In some embodiments, at least a portion of the paraffins and unreacted olefins stream is introduced directly into dehydrogenation-isomerization unit 114 via one or more entry points. Because the paraffins and unreacted olefins stream containing paraffins and unreacted olefins may be recycled to dehydrogenation-isomerization unit 114 as one stream, the process may be more efficient, resulting in an overall higher throughput. The higher throughput will increase the overall yield of the aliphatic alcohols.

In some embodiments, at least a portion of the paraffins and unreacted olefins stream may be introduced into hydrogenation unit 110 via fifth conduit 138. In certain embodiments, at least a portion of the paraffins and unreacted olefins stream may be combined with the first hydrocarbon stream in conduit 112 via fifth conduit 138. The combined stream may enter hydrogenation unit 110 and continue the process to produce aliphatic alcohols.

In some embodiments, an olefins and paraffins concentration in hydroformylation unit 120 may be adjusted depending on the source of the olefin stream entering the hydroformylation unit. A fourth hydrocarbon stream may be added upstream of hydroformylation unit 120 to produce a combined stream. In other embodiments, a fourth hydrocarbon stream may be introduced directly into hydroformylation unit 120 through one or more points. A fourth hydrocarbon stream may be introduced into third conduit 118 via sixth conduit 140 to produce a combined stream. The combined stream may enter

25
hydroformylation unit 120 via third conduit 118 to continue the process to produce aliphatic alcohols.

The fourth hydrocarbon stream may be from the same source as the first hydrocarbon stream. In some embodiments, a fourth hydrocarbon stream may be a hydrocarbon stream that includes olefins, paraffins, and/or hydrocarbon solvents derived from another source. The fourth hydrocarbon stream may include olefins and paraffins. In certain embodiments, an average carbon number of the hydrocarbons in the fourth hydrocarbon stream ranges from 7 to 18. In some embodiments, a paraffin content of the fourth hydrocarbon stream may be between about 60 percent and about 90 percent by weight. In other embodiments, a paraffin content of the fourth hydrocarbon stream may be greater than about 90 percent by weight.

In an embodiment, an olefin content of a fourth hydrocarbon stream ranges between about 1 percent and about 99 percent relative to the total hydrocarbon content. In certain embodiments, an olefin content of the fourth hydrocarbon stream may be between about 45 percent and about 99 percent by weight. In other embodiments, an olefin concentration of the fourth hydrocarbon stream may be greater than about 80 percent by weight.

In certain embodiments, dehydrogenation-isomerization unit 114 may be separated into a plurality of zones to control reaction temperatures and/or prevent unwanted side reactions (e.g., diene formation and/or cyclization reactions). A first hydrocarbon stream containing paraffins and unreacted olefins may be introduced into hydrogenation unit 110 via first conduit 112 as depicted for System 200 in FIG. 3A. Hydrogenation of olefins may occur as previously described in hydrogenation unit 110 to produce a second hydrocarbon stream. The second hydrocarbon stream may exit hydrogenation unit 110 via second conduit 116 and enter dehydrogenation-isomerization unit 114. Dehydrogenation-isomerization unit 114 may be divided into a plurality of zones. The plurality of zones may include, but is not limited to, a first reaction zone, a transition zone and a second reaction zone. In first reaction zone 210, at least a portion of the paraffins in the second hydrocarbon stream may be dehydrogenated to olefins to produce an olefinic stream. The process stream may then pass into second reaction zone 212. In second reaction zone 212, at least a portion of the olefins in the process stream may be isomerized to branched olefins to produce a third hydrocarbon stream.
In first reaction zone 210, the dehydrogenation catalyst may be selected from a wide range of catalyst types. For example, the catalyst may be based on a metal or metal compound deposited on a porous support. The metal or metal compound may be selected from, but is not limited to, chrome oxide, iron oxide and noble metals.


Reaction temperatures in first reaction zone 210 may range from about 300 °C to about 600 °C. In some embodiments, a reaction temperature in first reaction zone 210 may range from about 450 °C to about 550 °C. A total pressure in first reaction zone 210 may range from about 0.010 atmosphere (1 kPa) to about 25.0 atmospheres (2534 kPa). In certain embodiments, total pressure in first reaction zone 210 may range from about 0.010 atmospheres (1 kPa) to about 15.0 atmospheres (1520 kPa). In some embodiments, hydrogen may be fed together with the unreacted second hydrocarbon stream in order to prevent coking. Hydrogen and paraffins present in the unreacted second hydrocarbon stream may be fed at a hydrogen to paraffin molar ratio in a range from about 0.1 to about 20. In an embodiment, a hydrogen to paraffin molar ratio may be in a range of about 1 to about 10.

Residence time in first reaction zone 210 may be selected such that a conversion level of the paraffins to olefins is below about 50 mole percent. In certain embodiments, a conversion level of the paraffins to olefins may be kept in a range from about 10 mole percent to about 20 mole percent. By keeping the conversion level low, side reactions (e.g.,
diene formation and cyclization reactions) may be prevented. In some embodiments, an olefinic hydrocarbon stream may exit first reaction zone 210, pass through transition zone 214 and enter second reaction zone 212. Transition zone 214 may include heat exchanger 216. Heat exchanger 216 may reduce the temperature of the olefinic hydrocarbon stream.

In an embodiment, first reaction zone 210 and second reaction zone 212 in dehydrogenation-isomerization unit 114 may be separate units, as depicted in FIG. 3B, with heat exchanger 216 positioned between the two units.

After the olefinic hydrocarbon stream enters second reaction zone 212, at least a portion of the olefins are isomerized to branched olefins to produce a third hydrocarbon stream. The composition and level of branching of the third hydrocarbon stream may be performed by \(^1\)H NMR analysis. In an embodiment, an olefinic stream may exit first reaction zone 210 and directly enter second reaction zone 212 where at least a portion of the olefins in the olefinic stream are isomerized to branched olefins.

The catalyst used for isomerization of the olefins to branched olefins may be the same as described in U. S. Patent No. 5,648,584 to Murray, entitled “Process for Isomerizing Linear Olefins to Isoolefins” and U.S. Patent No. 5,648,585 to Murray et al., entitled “Process for Isomerizing Linear Olefins to Isoolefins.”

In an embodiment, linear olefins in a second hydrocarbon stream are isomerized in second reaction zone 212 by contacting at least a portion of the olefinic stream with a zeolite catalyst. The zeolite catalyst may have at least one channel with a crystallographic free channel diameter ranging from greater than 4.2 Å to less than about 7 Å. The zeolite catalyst may have an elliptical pore size large enough to permit entry of a linear olefin and at least partial diffusion of a branched olefin. The pore size of the zeolite catalyst may also be small enough to retard coke formation.

Temperatures in second reaction zone 212 may be from about 200 °C to about 500 °C to isomerize linear olefins to branched olefins. In some embodiments, reaction temperatures in the first reaction zone and the second reaction zone are substantially the same. In such embodiments, use of a heat exchanger is not required. Typically, however, the reaction temperature of second reaction zone 212 is less than the reaction temperature of the first reaction zone. The use of a heat exchanger lowers the temperature of the stream leaving the first reaction zone to the appropriate temperature for reaction in the second
reaction zone. Hydrocarbon partial pressure in the second reaction zone may be from about
0.1 atmosphere (10 kPa) to about 10 atmospheres (1013 kPa).

In some embodiments, the third hydrocarbon stream may exit a second reaction
zone and enter a separation unit. In the separation unit, branched olefins may be separated
from linear olefins and paraffins as previously described with regard to FIG. 1. Referring
to FIG. 3A, the third hydrocarbon stream may exit second reaction zone 212 via third
conduit 118 and enter hydroformylation unit 120. At least a portion of the olefins in the
third hydrocarbon stream may be hydroformylated to produce a hydroformylation reaction
stream as described for System 100 in FIG. 1. At least a portion of the hydroformylation
reaction stream may be separated into a bottom stream and a top stream using generally
known methods. The crude aliphatic alcohol product stream may be further purified as
described for System 100 to produce a paraffins and unreacted olefins stream and an
aliphatic alcohol product stream. The aliphatic alcohol product stream may include
branched aliphatic alcohols (e.g., branched primary aliphatic alcohols). The aliphatic
alcohol product stream may be recycled, transported to other processing units, sold, and/or
transported to storage vessels.

The hydroformylation reaction mixture stream may enter separator 130 via fourth
conduit 132. In separator 130, at least three streams, a bottom stream, a paraffins and
unreacted olefins stream and an aliphatic alcohol product stream, may be produced using
techniques previously described for System 100. At least a portion of the bottom stream
may be recycled to the hydroformylation unit via bottom stream recycle conduit 134. At
least a portion of the paraffins and unreacted olefins stream may be recycled, combined
with other process streams, transported to and/or stored on site. The aliphatic alcohol
product stream may be transported via product conduit 136 to be stored on site, sold
commercially, transported off-site and/or utilized in other processing units.

In an embodiment, at least a portion of the paraffins and unreacted olefins stream
may be combined with the second hydrocarbon stream to produce a combined hydrocarbon
stream via fifth conduit 138. The combined hydrocarbon stream may enter first reaction
zone 210 and undergo the dehydrogenation-isomerization process and hydroformylation
process to produce aliphatic alcohols. By recycling the paraffins and unreacted olefins
stream, the yield of product may be maximized. In an embodiment, a paraffins and
unreacted olefins stream may directly enter dehydrogenation-isomerization unit 114 through one or more points of entry.

In some embodiments, an olefins and paraffins concentration in hydroformylation unit 120 may be adjusted depending on the source of the olefin stream entering the hydroformylation unit as previously described for System 100. At least a portion of a fourth hydrocarbon stream may be introduced into third conduit 118 upstream of hydroformylation unit 120 via sixth conduit 140. The combined stream may be introduced into hydroformylation unit 120 via third conduit 118. At least a portion of the olefins in the combined stream may be hydroformylated to produce aliphatic alcohols. In an embodiment, a third hydrocarbon stream may be introduced directly into hydroformylation unit 120 through one or more points of entry.

A fourth hydrocarbon stream may be used to optimize the olefin concentration in hydroformylation unit 120 at a concentration sufficient to maximize hydroformylation of the olefin. In addition, the fourth hydrocarbon may optimize the ratio of linear to branched aliphatic groups in the aliphatic alcohol. The fourth hydrocarbon stream may be, but is not limited to, a hydrocarbon stream containing olefins, paraffins and/or hydrocarbon solvents.

In an embodiment, a catalyst in dehydrogenation-isomerization unit 114 may be used in a stacked bed configuration. A stacked bed configuration may allow for the use of one or more catalysts in the reactor. A catalyst for dehydrogenation of paraffins and a catalyst for isomerization of olefins may enhance the selectivity of the catalysts and/or the process. A stacked bed configuration of dehydrogenation-isomerization unit 114 is depicted for System 300 in FIG. 3. Operating conditions of the stacked bed configuration may be the same as for two-zone system described above for System 200. The second hydrocarbon stream may enter the dehydrogenation zone 310 via second conduit 116.

The dehydrogenation catalyst used in the stacked bed configuration may have nonacidic properties. The term "nonacidic," as used herein, refers to a catalyst that exhibits little skeletal isomerization activity. The dehydrogenation catalyst may include a noble metal, a Group TVA component, an alkali or alkaline earth component, a halogen component and/or a porous carrier material.

In certain embodiments, a noble metal may be dispersed throughout the dehydrogenation catalyst. An amount of noble metal may range between about 0.01 weight percent to about 5 weight percent, calculated on an elemental basis, of the final
dehydrogenation catalyst. In certain embodiments, a dehydrogenation catalyst includes about 0.1 weight percent to about 1 weight percent platinum. The noble metal may be incorporated into the catalytic composite by techniques known in the art (e.g., co-precipitation, co-gelation, ion exchange, impregnation, deposition from a vapor phase or from an atomic source) before incorporation of other catalytic components. In some embodiments, a noble metal may be incorporated into the catalytic composite during incorporation of other catalytic components. In other embodiments, a noble metal may be incorporated into the catalytic composite after incorporation of other catalytic components. In certain embodiments, a noble metal may be incorporated by impregnation of the carrier material with a solution or suspension of a decomposable compound of the noble metal. For example, platinum may be added to a catalytic support by commingling the platinum with an aqueous solution of chloroplatinic acid. In other embodiments, optional components (e.g., nitric acid) may be added to the impregnating solution to assist in dispersing or fixing the noble metal in the final catalyst composite.

The Group IVA component may include germanium, tin, lead or combinations thereof. In some embodiments, a Group IVA component may exist within the catalyst in an oxidation state above that of the noble metal. The Group IVA component may be present as an oxide. In certain embodiments, a Group IVA component may be combined with a carrier material. In some embodiments, a Group IVA component may be combined with the other catalytic components. In other embodiments, a Group IVA component may be dispersed throughout the catalyst. A Group IVA component may range between about 0.01 weight percent to about 5 weight percent, calculated on an elemental basis, of the final catalyst composite. In some embodiments, a catalyst includes about 0.2 weight percent to about 2 weight percent tin.

The Group IVA component may be incorporated in the catalytic composite according to generally known methods (e.g., co-precipitation, co-gelation, ion exchange and impregnation) before other catalytic components are incorporated. In some embodiments, a Group IVA component may be incorporated during incorporation of other catalytic components. In other embodiments, a Group IVA component may be incorporated after other catalytic components are incorporated. In some embodiments, a tin component may be incorporated by co-gelation with the porous carrier material. The tin may be incorporated in an alumina carrier material by mixing a soluble tin compound (e.g.,
stannous or stannic chloride) with an alumina hydrosol. A gelling agent (e.g., hexamethylenetetraamine) may be added to the tin-alumina hydrosol mixture. The tin-alumina hydrosol mixture may be dropped into an oil bath to form spheres containing alumina and tin. In other embodiments, a germanium component may be impregnated into a carrier material with a solution or suspension of a decomposable compound of germanium (e.g., germanium tetrachloride dissolved in an alcohol). In other embodiments, a lead component may be impregnated from a solution of lead nitrate in water.

In certain embodiments, an alkali or alkaline earth component may be included in the dehydrogenation catalyst. Alkali and alkaline earth component may include, but is not limited to, cesium, rubidium, potassium, sodium, lithium, barium, strontium, calcium and magnesium or mixtures thereof. The alkali or alkaline earth component may exist in the final catalytic composite in an oxidation state above that of the noble metal. The alkali or alkaline earth component may be present as an oxide. In some embodiments, an alkali or alkaline earth metal may be combined with the carrier material. In certain embodiments, an alkali or alkaline earth metal may be combined with other dehydrogenation catalytic components.

In other embodiments, an alkali or alkaline earth component may be dispersed throughout the catalytic composite. An amount of alkali or alkaline earth component may range from about 0.01 weight percent to 15 weight percent, calculated on an elemental basis, of the final catalytic composite. In other embodiments, a dehydrogenation catalyst includes about 1 weight percent to about 3 weight percent potassium. In certain embodiments, an atomic ratio of the alkali or alkaline earth component to the noble metal may be greater than at least about 10.

The alkali or alkaline earth component may be incorporated in the catalytic composite according to generally known methods (e.g., co-precipitation, co-gelation, ion exchange or impregnation) before other catalytic components are incorporated. In some embodiments, an alkali or alkaline earth component may be incorporated during incorporation of other catalytic components. In other embodiments, an alkali or alkaline earth component may be incorporated after other catalytic components are incorporated. For example, a potassium component may be impregnated into the carrier material with a solution of potassium nitrate. An atomic ratio of alkali or alkaline earth component to noble metal may be at least about 10. In certain embodiments, an atomic ratio of the alkali
or alkaline earth component to the noble metal component may range from about 15 to about 25.

A porous carrier material used in a dehydrogenation catalyst may include a porous, absorptive support with high surface area from about 25 m²/g to about 500 m²/g. The porous carrier material may have a melting point greater than the conditions utilized in the dehydrogenation zone. Examples of carrier materials include, but are not limited to, activated carbon, coke, charcoal, silica, silica gel, silicon carbide, synthetically prepared and/or naturally occurring clays and silicates, refractory inorganic oxides, crystalline zeolitic aluminosilicates, naturally occurring or synthetically prepared mordenite and/or faujasite, spinels or combinations of thereof. In certain embodiments, a carrier material may be gamma- or eta-alumina. In some embodiments, clays and silicates may or may not be acid treated (e.g., attapulgite, china clay, diatomaceous earth, fuller's earth, kaolin, kieselguhr, ceramics, porcelain, crushed firebrick, bauxite). Examples of refractory inorganic oxides include alumina, titanium dioxide, zirconium dioxide, chromium oxide, beryllium oxide, vanadium oxide, cerium oxide, hafnium oxide, zinc oxide, magnesia, boria, thorium, silica-alumina, silica-magnesia, chromia-alumina, alumina-boria, and silica-zirconia. Zeolitic aluminosilicates may be, in some embodiments, in the hydrogen form. In other embodiments, zeolitic aluminosilicates may be in a form that may be exchanged with metal cations. Examples of spinels include, but are not limited to, MgAl₂O₄, FeAl₂O₄, ZnAl₂O₄, CaAl₂O₄, and other like compounds having the formula MO-Al₂O₃ in which M is a metal having a valence of 2.

In certain embodiments, an alumina carrier material used in a dehydrogenation catalyst may be prepared in any suitable manner from synthetic or naturally occurring raw materials. The alumina carrier may be formed in any desired shape (e.g., spheres, pills, cakes, extrudates, powders, granules). The alumina carrier may be utilized in any particle size. In certain embodiments, a sphere shape may be utilized. The particle may be about 1/16 inch in diameter. In certain embodiments, a particle diameter of less than about 1/32 inch may be utilized.

Alumina spheres may be prepared, in some embodiments, by converting aluminum metal into an alumina sol. An alumina sol may be prepared by reacting aluminum metal with a suitable peptizing acid and water. The resulting alumina sol and a gelling agent may be dropped into an oil bath to form spherical particles of alumina gel. The resulting
alumina gel may be converted to gamma- or eta-alumina carrier material using known techniques (e.g., by aging, drying and calcining).

In other embodiments, alumina cylinders may be prepared by mulling alumina powder with water and a suitable peptizing agent (e.g., nitric acid) to form an extrudable composition. The composition may be extruded through a suitably sized die then cut to form extrudate particles. Other shapes of the alumina carrier material may be prepared by conventional methods. After the alumina particles are shaped, they may be dried and calcined. The alumina carrier may be subjected to intermediate treatments (e.g., washing with water or a solution of ammonium hydroxide) during preparation.

The dehydrogenation catalyst may include a halogen component. The halogen component may include, but is not limited to, fluorine, chlorine, bromine, iodine or mixtures thereof. The halogen component may be present in a combined state with the porous carrier material. In certain embodiments, a halogen component may be dispersed throughout the catalytic composite. A halogen component may range from at least about 0.2 weight percent to about 15 weight percent, calculated on an elemental basis, of the final catalytic composite. In certain embodiments, a dehydrogenation catalyst contains about 1 weight percent to about 3 weight percent chlorine.

In certain embodiments, a catalyst composition may include at least about 0.2 weight percent, calculated on an elemental basis, of a halogen component. The halogen component in the catalyst may improve the activity of the catalyst for dehydrogenating hydrocarbons. In some embodiments, an active halogen component may suppress carbon formation on the catalyst during the dehydrogenation process. An advantage of the catalyst composition may be that undesirable isomerization or cracking side reactions may be inhibited. In certain embodiments, halogen content may increase the acidity of the catalyst.

The acidity may be lowered by steaming the dehydrogenation catalyst to remove excess halogen from the dehydrogenation catalyst.

A halogen component may be incorporated in the catalytic composite in any suitable manner. The incorporation of the halogen may be before preparation of the carrier material. In some embodiments, incorporation of a halogen may be during incorporation of other catalytic components. In other embodiments, incorporation of a halogen may be after other catalytic components are incorporated. In certain embodiments, an alumina sol carrier may contain a halogen, which may contribute to at least some portion of the halogen
content in the final catalyst composite. In some embodiments, a halogen component, or a portion thereof, may be added to the catalyst composite during the incorporation of the carrier material with other catalyst components (e.g., using chloroplatinic acid to impregnate the platinum component). In other embodiments, a halogen component or a portion thereof may be added to the catalyst composite by contacting the catalyst with the halogen. In some embodiments, a halogen may be added to the catalyst as a compound, solution, suspension or dispersion containing the halogen, (e.g., hydrochloric acid) before or after other catalyst components are incorporated with the carrier material. In certain embodiments, a halogen component or a portion thereof may be incorporated by contacting the catalyst with a compound, solution, suspension or dispersion containing the halogen in a subsequent catalyst regeneration step. In the regeneration step, carbon deposited on the catalyst as coke during use of the catalyst in a hydrocarbon conversion process may be burned off the catalyst. The noble metal on the catalyst may be redistributed to provide a regenerated catalyst with performance characteristics similar to those of the fresh catalyst.

The halogen component may be added during the carbon burn step or during the noble metal redistribution step (e.g., contacting the catalyst with a hydrogen chloride gas). In some embodiments, a halogen component may be added to the catalyst composite by adding the halogen or a compound, solution, suspension or dispersion containing the halogen (e.g., propylene dichloride) to the hydrocarbon feed stream. In other embodiments, a halogen component may be added to the recycle gas during operation of the dehydrogenation unit.

In some embodiments, a dehydrogenation catalyst may include a sulfur component ranging from about 0.01 weight percent to about 10 weight percent, calculated on an elemental basis, of the final catalytic composition. The sulfur component may be incorporated into the catalytic composite in any suitable manner. In certain embodiments, sulfur or a compound containing sulfur (e.g., hydrogen sulfide or a lower molecular weight mercaptan) may be contacted with the catalyst composition in the presence of hydrogen at a temperature ranging from about 10 °C to about 540 °C under anhydrous conditions. A hydrogen to sulfur ratio, in some embodiments, may be about 100.

The dehydrogenation catalyst, in some embodiments, may also contain other, additional components or mixtures thereof, which act alone or in concert, as catalyst modifiers to improve catalyst activity, selectivity or stability. Examples of catalyst
modifiers include, but are not limited to, antimony, arsenic, beryllium, bismuth, cadmium, calcium, chromium, cobalt, copper, gallium, gold, indium, iron, lithium, manganese, molybdenum, nickel, rhenium, scandium, silver, tantalum, thallium, titanium, tungsten, uranium, zinc and zirconium. Catalytic modifiers may be added in any suitable manner to the carrier material during preparation of the dehydrogenation catalyst. In other embodiments, catalytic modifiers may be added in any suitable manner after preparation of the dehydrogenation catalyst. In some embodiments, catalytic modifiers may be added in any suitable manner to the catalytic composite before other catalytic components are incorporated. In certain embodiments, catalytic modifiers may be added during incorporation of other catalytic components. In other embodiments, catalytic modifiers may be added after other catalytic components are incorporated. A description of a dehydrogenation catalyst may be found in U.S. Patent No. 4,506,032 to Imai et al., entitled “Dehydrogenation Catalyst Composition.”

The olefinic hydrocarbon stream may pass into isomerization zone 312. In certain embodiments, a temperature decrease from dehydrogenation zone 310 to isomerization zone 312 may be necessary to prevent cracking of the olefinic hydrocarbon stream as it enters the isomerization zone. Cool hydrogen gas may be introduced to dehydrogenation zone 310 via gas conduit 314 to control temperatures in dehydrogenation zone 310. In isomerization zone 312, at least a portion of the olefins in the olefinic hydrocarbon stream may be isomerized to branched olefins to produce a third hydrocarbon stream.

In certain embodiments, an isomerization catalyst may be the same as described for isomerization of olefins in System 200. A description of the isomerization catalyst may be found in U.S. Patent No. 5,510,306 to Murray, entitled “Process For Isomerizing Linear Olefins to Isoolefins.” In some embodiments, about 0.01 weight percent to about 5 weight percent of a noble metal may be added to an isomerization catalyst used in a stacked bed configuration to increase the dehydrogenation activity of the zeolitic catalyst. Common metal incorporation methods (e.g., impregnation, noble metal ion exchange, co-mulling) may be used to incorporate a noble metal (e.g., platinum, palladium) into a zeolite to produce a working catalyst useful in the dehydrogenation-isomerization of paraffins.

The third hydrocarbon stream may exit isomerization zone 312 and enter hydroformylation unit 120 via third conduit 118. At least a portion of the olefins in the third hydrocarbon stream may be hydroformylated to produce a hydroformylation reaction.
stream as described for System 100. At least a portion of the hydroformylation reaction stream may be separated into a bottoms stream and a top stream using generally known methods. The top stream may be purified and separated as described for System 100 to produce a paraffins and unreacted olefins stream and a crude aliphatic alcohol product stream. The crude aliphatic alcohol product stream may be further purified as described for System 100 to produce an aliphatic alcohol product stream. The hydroformylation reaction mixture stream may enter separator 130 via fourth conduit 132. In separator 130 at least two streams, a bottom stream and top stream, may be produced as previously described for System 100. The bottom stream may be recycled to hydroformylation unit 120 via bottom stream recycle conduit 134. The top stream may be purified and separated into at least two streams, a paraffins and unreacted olefins stream and a crude aliphatic alcohol product stream. At least a portion of the paraffins and unreacted olefins stream may be recycled, combined with other process streams, sent to other processing units and/or sent to a storage vessel. The crude aliphatic alcohol product stream may be further purified as described for System 100 to produce an aliphatic alcohol product stream. The aliphatic alcohol product stream may include branched aliphatic alcohols (e.g., branched primary aliphatic alcohols). The aliphatic alcohol product stream may be transported via product conduit 136 to be stored on site, sold commercially, transported off-site and/or utilized in other processing units.

At least a portion of the paraffins and unreacted olefins stream may be combined with the second hydrocarbon stream in second conduit 116 to produce a combined hydrocarbon stream via fifth conduit 138. The combined hydrocarbon stream may enter dehydrogenation zone 310 of dehydrogenation-isomerization unit 114 via second conduit 116. The combined hydrocarbon stream entering dehydrogenation zone 310 continues the dehydrogenation-isomerization process and hydroformylation process to produce aliphatic alcohols. By recycling the paraffins and unreacted olefins stream, the yield of product may be maximized. In an embodiment, a paraffins and unreacted olefins stream may directly enter dehydrogenation-isomerization unit 114 through one or more entry points.

In some embodiments, an olefin and paraffin concentration in hydroformylation unit 120 may be adjusted depending on the source of the olefin stream entering the hydroformylation unit. At least a portion of a fourth hydrocarbon stream may be introduced into third conduit 118 upstream of the hydroformylation unit as previously
described for System 100. The combined stream may be introduced into hydroformylation unit 120 via third conduit 118. At least a portion of the olefins in the combined stream may be hydroformylated to produce aliphatic alcohols.

Aliphatic alcohols produced in Systems 100, 200 and 300 may be converted to oxy alcohols, sulfates or other commercial products. At least a portion of the aliphatic alcohols in the alcohol product stream may be reacted in an oxyalkylation unit with an epoxide (e.g., ethylene oxide, propylene oxide, butylene oxide) in the presence of a base to produce an oxyalkyl alcohol. Condensation of an alcohol with an epoxide allows the alcohol functionality to be expanded by one or more oxy groups. The number of oxy groups may range from 3 to 12. For example, reaction of an alcohol with ethylene oxide may produce alcohol products having between 3 to 12 ethoxy groups. Reaction of an alcohol with ethylene oxide and propylene oxide may produce alcohols with an ethoxy/proproxy ratio of ethoxy to proproxy groups from about 4:1 to about 12:1. In some embodiments, a substantial proportion of alcohols moieties may become combined with more than three ethylene oxide moieties. In other embodiments, an approximately equal proportion may be combined with less than three ethylene oxide moieties. In a typical oxyalkylation product mixture, a minor proportion of unreacted alcohol may be present in the product mixture. In an embodiment, at least a portion of the aliphatic alcohol product stream may be formed by condensing a C₅ to C₅¹ aliphatic alcohol with an epoxide. In certain embodiments, a C₅ to C₁₅ branched primary alcohol may be condensed with ethylene oxide and/or propylene oxide. In other embodiments, a C₁₁ to C₁₇ branched primary alcohol may be condensed with ethylene oxide and/or propylene oxide. The resulting oxyalkyl alcohols may be sold commercially, transported off-site, stored on site and/or used in other processing units. In some embodiments, an oxyalkyl alcohol may be sulfated to form an anionic surfactant.

In an embodiment, at least a portion of the alcohols in the aliphatic alcohol product stream may be added to a base. The base may be an alkali metal or alkaline earth metal hydroxide (e.g., sodium hydroxide or potassium hydroxide). The base may act as a catalyst for the oxyalkylation reaction. An amount from about 0.1 percent by weight to about 0.6 percent by weight of a base, based on the total weight of alcohol, may be used for oxyalkylation of an alcohol. In an embodiment, a weight percent of a base may range from about 0.1 percent by weight to 0.4 percent by weight based on the total alcohol amount. The reaction of the alcohol with the base may result in formation of an alkoxide. The
resulting alkoxide may be dried to remove any water present. The dried alkoxide may be reacted with an epoxide. An amount of epoxide used may be from about 1 mole to about 12 moles of epoxide per mole of alkoxide. A resulting alkoxide-epoxide mixture may be allowed to react until the epoxide is consumed. A decrease in overall reaction pressure may indicate that the reaction is complete.

Reaction temperatures in an oxyalkylation unit may range from about 120 °C to about 220 °C. In an embodiment, reaction temperatures may range from about 140 °C to about 160 °C. Reaction pressures may be achieved by introducing to the reaction vessel the required amount of epoxide. Epoxides have a high vapor pressure at the desired reaction temperature. For consideration of process safety, the partial pressure of the epoxide reactant may be limited, for example, to less than about 4 atmospheres (413 kPa). Other safety measures may include diluting the reactant with an inert gas such as nitrogen. For example, inert gas dilution may result in a vapor phase concentration of reactant of about 50 percent or less. In some embodiments, an alcohol-epoxide reaction may be safely accomplished at a greater epoxide concentration, a greater total pressure and a greater partial pressure of epoxide if suitable, generally known, safety precautions are taken to manage the risks of explosion. With respect to ethylene oxide, a total pressure from about 3 atmospheres (304 kPa) to about 7 atmospheres (709 kPa) may be used. Total pressures of ethylene oxide from about 1 atmosphere (101 kPa) to about 4 atmospheres (415 kPa) may be used in certain embodiments. In an embodiment, total pressures from about 1.5 atmospheres (150 kPa) to about 3 atmospheres (304 kPa) with respect to ethylene oxide may be used. The pressure may serve as a measure of the degree of the reaction. The reaction may be considered substantially complete when the pressure no longer decreases with time.

Aliphatic alcohols and oxyalkyl alcohols may be derivatized to form compositions (e.g., sulfonates, sulfates, phosphates) useful in commercial product formulations (e.g., detergents, surfactants, oil additives, lubricating oil formulations). For example, alcohols may be sulfurized with SO₂ to produce sulfates. The term “sulfurized” refers to a sulfur atom or sulfur containing functionality being added to a carbon or oxygen. Sulfurization processes are described in U.S. Patent No. 6,462,215 to Jacobson et al., entitled “Sulfonation, Sulfation and Sulfamation”; U.S. Patent No. 6,448,435 to Jacobson et al., entitled “Sulfonation, Sulfation and Sulfamation”; U.S. Patent No. 3,462,525 to Levinsky

A general class of aliphatic alcohol sulfates may be characterized by the chemical formula: \((R-O-(A)_x-SO_3)_nM\). \(R'\) represents the aliphatic moiety. \(A\) represents a moiety of an alkylene oxide; \(x\) represents the average number of \(A\) moieties per R-O moiety and may range from 0 to 15; and \(n\) is a number depending on the valence of cation \(M\).

Examples of cation \(M\) include, but are not limited to, alkali metal ions, alkaline earth metal ions, ammonium ions and/or mixtures thereof. Examples of cations include, but are not limited to, magnesium, potassium, monoethanol amine, diethanol amine or triethanol amine.

Aliphatic and oxyalkyl alcohols may be sulfated in a sulfation unit. Sulfation procedures may include the reaction of sulfur trioxide (SO₃), chlorosulfonic acid (ClSO₂H), sulfamic acid (NH₂SO₂H) or sulfuric acid with an alcohol. In an embodiment, sulfur trioxide in concentrated (e.g., fuming) sulfuric acid may be used to sulfate alcohols. The concentrated sulfuric acid may have a concentration of about 75 percent by weight to about 100 percent by weight in water. In an embodiment, concentrated sulfuric acid may have a concentration of about 85 percent by weight to about 98 percent by weight in water. The amount of sulfur trioxide may range from about 0.3 mole to about 1.3 moles of sulfur trioxide per mole of alcohol. In certain embodiments, an amount of sulfur trioxide may range from about 0.4 moles to about 1.0 moles of sulfur trioxide per mole of alcohol.

In an embodiment, a sulfur trioxide sulfation procedure may include contacting a liquid alcohol or an oxyalkyl alcohol and gaseous sulfur trioxide in a falling film sulfator to produce a sulfuric acid ester of the alcohol. The reaction zone of the falling film sulfator may be operated at about atmospheric pressure and at a temperature in the range from about 25 °C to about 70 °C. The sulfuric acid ester of the alcohol may exit the falling film sulfator and enter a neutralization reactor. The sulfuric acid ester may be neutralized with an alkali metal solution to form the alkyl sulfate salt or the oxyalkyl sulfate salt. Examples of an alkali metal solution may include solutions of sodium or potassium hydroxide.
The derivatized alcohols may be used in a wide variety of applications. An example of an application includes detergent formulations. Detergent formulations include, but are not limited to, granular laundry detergent formulation, liquid laundry detergent formulations, liquid dishwashing detergent formulations and miscellaneous formulations. Examples of miscellaneous formulations may include general purpose cleaning agents, liquid soaps, shampoos and liquid scouring agents.

Granular laundry detergent formulations may include a number of components besides the derivatized alcohols (e.g., surfactants, builders, co-builders, bleaching agents, bleaching agent activators, foam controlling agents, enzymes, anti-graying agents, optical brighteners and stabilizers). Examples of other surfactants may include ionic, nonionic, amphoteric or cationic surfactants.

Liquid laundry detergent formulations may include the same components as granular laundry detergent formulations. In certain embodiments, liquid laundry detergent formulations may include less of an inorganic builder component than granular laundry detergent formulations. Hydrogropes may be present in the liquid detergent formulations. General purpose cleaning agents may include other surfactants, builders, foam control agents, hydrogropes and solubilizer alcohols.

Examples

Example 1: ISOMERIZATION OF OLEFINs IN A FISCHER-TROPSCH DERIVED HYDROCARBON STREAM. Carbon monoxide and hydrogen were reacted under Fischer-Tropsch process conditions to yield a hydrocarbon mixture of linear paraffins, linear olefins, a minor amount of dienes and a minor amount of oxygenates. The Fischer-Tropsch hydrocarbon stream was separated into different hydrocarbon streams using fractional distillation techniques. A hydrocarbon stream containing olefins and paraffins with an average number of carbon atoms from 8 to 10 was obtained. The composition of the resulting C₈-C₁₀ hydrocarbon stream was analysed by gas chromatography and is tabulated in Table 1.
A zeolite catalyst used for isomerization of linear olefins in the hydrocarbon stream was prepared in the following manner. Ammonium-ferrierite (645 grams) exhibiting a 5.4% loss on ignition and exhibiting the following properties: molar silica to alumina ratio of 62:1, surface area of 369 square meters per gram (P/Po=0.03), soda content of 480 ppm and n-hexane sorption capacity of 7.3 g per 100 g of ammonium-ferrierite was loaded into a Lancaster mix muller. CATAPAL® D alumina (91 grams) exhibiting a loss on ignition of 25.7% was added to the muller. During a five-minute mulling period, 152 milliliters of deionized water was added to the alumina/ammonium-ferrierite mixture. Next, a mixture of 6.8 grams glacial acetic acid, 7.0 grams of citric acid and 152 milliliters of deionized water was slowly added to the alumina/ammonium-ferrierite mixture in the muller to peptize the alumina. The resulting alumina/ammonium-ferrierite/acid mixture was mulled for 10 minutes. Over a period of 15 minutes, a mixture of 0.20 grams of tetramine palladium nitrate in 153 grams of deionized water was slowly added to mulled alumina/ammonium-ferrierite/acid mixture. The resulting mixture exhibited a 90:10 ratio of zeolite to alumina and a loss on ignition of 43.5%. The zeolite/alumina mixture was shaped by extruding the mixture through a stainless steel die plate (1/16” holes) of a 2.25 inch Bonnot extruder.

The moist zeolite/alumina extrudate was dried at 125°C for 16 hours. After drying, the zeolite/alumina extrudate was longsbroked manually. The zeolite/alumina extrudate
was calcined in flowing air at 200°C for two hours. The temperature was raised to a maximum temperature of 500°C and the zeolite/alumina extrudate was calcined for an additional two hours to yield an isomerization catalyst. The isomerization catalyst was allowed to cool in a desiccator under a nitrogen atmosphere.

Stainless steel tubing, 1 inch OD, 0.6 inch ID and 26 inches long, was used as an isomerization reactor. A thermowell extended 20 inches from the top of the stainless steel reactor tube. To load the reactor tube, the reactor tube was inverted and a piece of glass wool was transferred down the wall of the reactor tube, over the thermowell and positioned at the bottom of the reactor tube to serve as a plug for the reactor tube. Silicon carbide (20 mesh) was added to a depth of about 6 inches to the reactor tube. A second piece of glass wool was placed over the silicon carbide. A mixture of 6.0 grams of the isomerization catalyst particles (6-20 mesh) and 45 grams of fresh silicon carbide (60-80 mesh) was added to the reactor tube in two parts. The two-part addition distributed the isomerization catalyst evenly in the reactor tube and resulted in an isomerization catalyst bed of about 10 inches in length. A third piece of glass wool was added to the top of the catalyst in the reactor tube. Silicon carbide (20 mesh) was layered onto the third piece of glass wool. A fourth piece of glass wool was positioned over the silicon carbide to serve as a plug for the bottom of the reactor tube. To monitor the temperature of the reaction at various points in the reactor tube, a multipoint thermocouple was inserted into the thermowell of the reactor tube. The temperature above, below and at three different places in the catalyst bed was monitored. The reactor tube was inverted and installed in the furnace. The reactor tube was heated to the operating temperature of 280 °C over a four-hour period under flowing nitrogen. Once the temperature of 280 °C was obtained, the reactor tube was held at the operating temperature for an additional two hours to condition the isomerization catalyst.

After conditioning the isomerization catalyst, the hydrocarbon stream was pumped through the reactor tube at a flow rate of 60 g/hr. Nitrogen, at a flow rate of 6 L/hr, was passed over the isomerization catalyst simultaneously with the hydrocarbon stream. The hydrocarbon stream was vaporized before contacting the isomerization catalyst. The reactor tube was operated at an outlet pressure of 20 kPa above atmospheric pressure.

In Table 2, the weight percent of C₆-C₁₀ branched olefins, C₆-C₁₀ linear olefins and C₆-C₁₀ paraffins in the hydrocarbon stream at 0 hours and in the reactor tube effluent after 24 and 48 hours of isomerization is tabulated. Greater than 90% of the linear olefins in the
hydrocarbon stream were converted into branched olefins in the isomerization reactor. During the isomerization step, a small amount of material boiling below C₈ was generated from cracking side reactions. In addition, a portion of the C₉-C₁₁ alcohols present in the feed was dehydrated to yield additional olefins in the product. The average number of alkyl branches on the C₈-C₁₀ olefins in the product was found to be 1.0 as determined by ¹H NMR analysis.

<table>
<thead>
<tr>
<th>Fischer-Tropsch Hydrocarbon Stream Composition During Isomerization Reaction</th>
<th>0 Hr Wt.%</th>
<th>24 Hr Wt.%</th>
<th>48 Hr Wt.%</th>
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<tr>
<td>C₈-C₁₀ branched olefins</td>
<td>0.46</td>
<td>33.04</td>
<td>33.16</td>
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<tr>
<td>C₈-C₁₀ linear olefins</td>
<td>32.19</td>
<td>2.52</td>
<td>2.54</td>
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<td>C₈-C₁₀ paraffins</td>
<td>63.19</td>
<td>63.32</td>
<td>63.27</td>
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<tr>
<td>Branched to linear C₈-C₁₀ olefin ratio</td>
<td>0.1</td>
<td>13.1</td>
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</tbody>
</table>

Example 2. ISOMERIZATION OF 1-DODECENE. 1-Dodecene was obtained from Shell Chemical Co. The composition of 1-dodecene, as assayed by gas chromatography, is tabulated in Table 3.

<table>
<thead>
<tr>
<th>1-Dodecene Composition</th>
<th>Wt.%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-Dodecene</td>
<td>98.0</td>
</tr>
<tr>
<td>Other C₁₀-C₁₄ olefins</td>
<td>1.2</td>
</tr>
<tr>
<td>&lt;C₁₀ hydrocarbons</td>
<td>0.2</td>
</tr>
<tr>
<td>&gt;C₁₄ hydrocarbons</td>
<td>0.2</td>
</tr>
<tr>
<td>Paraffins</td>
<td>0.4</td>
</tr>
<tr>
<td>Total C₁₀-C₁₄ hydrocarbons</td>
<td>99.6</td>
</tr>
</tbody>
</table>

1-dodecene was isomerized using the same reactor tube design and isomerization catalyst preparation as described in Example 1. A stream of 1-dodecene was pumped through a reactor tube at a flow rate of 90 g/hr. Nitrogen, at a flow rate of 6 L/hr, was passed over the isomerization catalyst simultaneously with the stream of 1-dodecene. The stream of 1-dodecene was vaporised before contacting the isomerization catalyst. The reactor tube was operated at an outlet pressure of 20 kPa above atmospheric pressure and a temperature of 290°C.
Table 4 is a tabulation of the weight percent of less than C_{10}, C_{10}-C_{14} and greater than C_{14} molecules in 1-dodecene at 0 hours and the reactor tube effluent after 168 and 849 hours. Linear C_{10}-C_{14} olefins were converted in a 94% yield to branched C_{10}-C_{14} olefins after a 168 hr processing time. During the isomerization step, less than 3 weight percent of material boiling below C_{10} was generated from cracking side reactions. The average number of alkyl branches on the C_{10}-C_{14} olefins in the product was determined to be 1.3 by $^1$H NMR analysis.

<table>
<thead>
<tr>
<th>1-Dodecene Stream Composition During Isomerization Reaction</th>
<th>0 Hr Wt.%</th>
<th>168 Hr Wt.%</th>
<th>849 Hr Wt.%</th>
</tr>
</thead>
<tbody>
<tr>
<td>$&lt;$C_{10} hydrocarbons</td>
<td>0.2</td>
<td>2.5</td>
<td>2.4</td>
</tr>
<tr>
<td>C_{10}-C_{14} hydrocarbons</td>
<td>99.6</td>
<td>97.2</td>
<td>97.4</td>
</tr>
<tr>
<td>$&gt;$C_{14} hydrocarbons</td>
<td>0.2</td>
<td>0.3</td>
<td>0.2</td>
</tr>
<tr>
<td>Branched C_{10}-C_{14} olefins</td>
<td>0.6</td>
<td>93.2</td>
<td>93.4</td>
</tr>
<tr>
<td>Linear C_{10}-C_{14} olefins</td>
<td>99.0</td>
<td>2.8</td>
<td>2.9</td>
</tr>
<tr>
<td>Paraffins</td>
<td>1.0</td>
<td>2.0</td>
<td>1.9</td>
</tr>
</tbody>
</table>

EXAMPLE 3. DEHYDROGENATION OF DODECANE WITH MINIMAL ISOMERIZATION. Dodecane was obtained from Aldrich Chemical Company and stored under nitrogen before being processed. The composition of dodecane, as assayed by gas chromatography, is tabulated in Table 5.

<table>
<thead>
<tr>
<th>Dodecane Composition</th>
<th>Wt.%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dodecane</td>
<td>99.3</td>
</tr>
<tr>
<td>$&lt;$C_{10} hydrocarbons</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>C_{10}, C_{11}, C_{13} and C_{14} hydrocarbons</td>
<td>&lt;0.6</td>
</tr>
<tr>
<td>$&gt;$C_{14} hydrocarbons</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>Other C_{10}-C_{14} olefins</td>
<td>&lt;0.1</td>
</tr>
</tbody>
</table>

A paraffin dehydrogenation catalyst was prepared according to Example 1 (catalyst A) of U.S. Patent No. 4,430,517 to Imai et al., entitled “Dehydrogenation Process Using A Catalytic Composition.” The resulting catalyst included 0.8 wt.% platinum, 0.5 wt.% tin, 2.7 wt.% tin, 2.7 wt.% potassium and 1.3 wt.% chlorine on a gamma-alumina support. The atomic ratio of potassium to platinum for this catalyst was 16.8.
The dehydrogenation catalyst was prepared by dissolving substantially pure aluminum pellets in a hydrochloric acid solution. An amount of stannic chloride was added to the resulting solution to provide a final composite containing 0.5 weight % tin and stirred to distribute the tin component evenly throughout the mixture.

Hexamethylenetetramine was added to the resulting tin mixture and the resulting tin-amine mixture was dropped into an oil bath in a manner to form spherical particles having an average particle diameter of about 1/16 inch. The spheres were aged, washed with an ammoniacal solution, dried and calcined to form a spherical gamma-alumina carrier material. The resulting spheres contained about 0.5 weight % tin in the form of tin oxide.

More details about the method of preparing the alumina carrier material are disclosed in U.S. Patent No. 2,620,314 to Hoeskstra, entitled, “Spheroidal Alumina.”

The tin-alumina composite was contacted with a deionized solution of chloroplatinic acid and hydrochloric acid (2 weight percent based on alumina weight) in a rotary drier for 15 minutes at room temperature. The amount of chloroplatinic acid used was the amount necessary to incorporate 0.8 weight percent platinum into the tin-alumina composite. The solution was then heated and purged with nitrogen to remove water resulting in a platinum-chlorine-tin-alumina composite. The incorporated chlorine was removed by heating the platinum-chlorine-tin-alumina composite to 550 °C and treating the composite with a 50/50 air/80 °C steam mixture at a gas hourly space velocity (GHSV) of 300 hr⁻¹. After treatment with the air/steam mixture, the platinum-tin-alumina composite contained less than 0.1 weight percent chlorine.

The platinum-tin-alumina composite was contacted with a deionized water solution of potassium nitrate. The amount of potassium nitrate used was the amount necessary to incorporate 2.7 weight percent of potassium in the platinum-tin-alumina composite. The water was removed from the platinum-tin-potassium-alumina composite by heating the composite to 100 °C under a purge of dry air (1000 hr⁻¹ GHSV) for 0.5 hour. The temperature was raised to 525 °C and the platinum-tin-potassium alumina composite was treated with a stream of hydrochloric acid (12 cc/hr, 0.9 M HCl) and a stream of 50/50 air/80 °C steam mixture (300 hr⁻¹ GHSV) to incorporate chlorine into the platinum-tin-potassium-alumina composite. The platinum-tin-potassium-chlorine-alumina composite was dried at 525 °C under a purge of dry air (1000 hr⁻¹ GHSV). The resulting catalyst
spheres had an average particle diameter of 1/16 inch and were crushed and sized into 6-20 mesh particle before testing.

Stainless steel tubing, 1 inch OD, 0.6 inch ID and 26 inches long, was used as an isomerization reactor. A thermowell extended 20 inches from the top of the stainless steel reactor tube. To load the reactor tube, the reactor tube was inverted and a piece of glass wool was transferred down the wall of the reactor tube, over the thermowell and positioned at the bottom of the reactor tube to serve as a plug for the reactor tube. Silicon carbide (20 mesh) was added to a depth of about 6 inches to the reactor tube. A second piece of glass wool was placed over the silicon carbide. A mixture of 6.0 grams of platinum-tin on alumina catalyst particles (6-20 mesh) and 45 grams of fresh silicon carbide (60-80 mesh) was added to the reactor tube in two parts. The two-part addition distributed the catalyst evenly in the reactor tube and resulted in a catalyst bed of about 10 inches in length. A third piece of glass wool was added to the top of the catalyst in the reactor tube. Silicon carbide (20 mesh) was layered onto the third piece of glass wool. A fourth piece of glass wool was positioned over the silicon carbide to serve as a plug for the bottom of the reactor tube. To monitor the temperature of the reaction at various points in the reactor tube, a multipoint thermocouple was inserted into the thermowell of the reactor tube. The temperature above, below and at three different places in the catalyst bed was monitored. The reactor tube was inverted and installed in the furnace. The reactor tube was purged with nitrogen. The reactor tube was heated to the operating temperature of 425°C over a four-hour period under flowing nitrogen (250 standard liters per hour). Once the temperature of 425°C was obtained, the reactor tube was held at the operating temperature for an additional two hours. The catalyst was presulfided by flowing a 1% mixture of hydrogen sulfide gas in hydrogen gas at 425 °C for five minutes through the reactor tube.

After 5 minutes, the hydrogen sulfide in hydrogen gas flow was switched to a hydrogen gas flow through the reactor tube.

After presulfiding the catalyst, the reactor tube was maintained at 425 °C for eight hours. After eight hours, the reactor tube pressure was increase to 25 psig with hydrogen gas. Dodecane was pumped through the reactor tube at a flow rate of 40 g/hr at a hydrogen flow rate of 125 standard liters per hour. After four hours, the dodecane stream was increased to 80 g/hr. After obtaining a flow rate of 80 g/hr, the reactor tube temperature
was raised to 460 °C. The reactor tube was sampled every eight hours after obtaining the operating temperature of 460°C.

After twenty-four hours the weight percent of dodecane was 11.4 weight percent as depicted in Table 6. At a temperature of 479 °C, the conversion of dodecane to olefins was 16 weight percent after twenty-four hours. Of the olefins, formed 84 weight percent were mono olefins, 4.1 weight percent were aromatic compounds and 7.5 weight percent were di-olefins. Of the total amount of olefins formed, 6 percent were branched, as determined by $^1$H NMR analysis.

<table>
<thead>
<tr>
<th>Test Results</th>
<th></th>
</tr>
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<tbody>
<tr>
<td>Conversion (wt.%) after 24 hours on-stream at 460 °C.</td>
<td>11.4</td>
</tr>
<tr>
<td>Temperature required for 16 wt. % conversion</td>
<td>479 °C</td>
</tr>
<tr>
<td>Selectivity to mono olefins at 16 wt. % conversion</td>
<td>84 wt.%</td>
</tr>
<tr>
<td>Selectivity to aromatics at 16 wt. % conversion</td>
<td>4.1 wt.%</td>
</tr>
<tr>
<td>Selectivity to di-olefins at 16 wt. % conversion</td>
<td>7.5 wt.%</td>
</tr>
<tr>
<td>% Branched C$<em>{12}$ olefins in total C$</em>{12}$ olefins, (wt.%)</td>
<td>6</td>
</tr>
</tbody>
</table>

**EXAMPLE 4: DEHYDROGENATION-ISOMERIZATION OF DODECANE:** Dodecane was obtained from Aldrich Chemical Company and stored under nitrogen before being processed. The composition of dodecane, as assayed by gas chromatography, is tabulated in Table 5.

A dehydrogenation-isomerization catalyst was prepared in the following manner. Ammonium-ferrierite (645 grams) exhibiting a 5.4% loss on ignition and exhibiting the following properties: molar silica to alumina ratio of 62:1, surface area of 369 square meters per gram (P/Po=0.03), soda content of 480 ppm and n-hexane sorption capacity of 7.3 g per 100 g of ammonium-ferrierite was loaded into a Lancaster mix muller.

CATAPAL® D alumina (91 grams) exhibiting a loss on ignition of 25.7% was added to the muller. During a five-minute mulling period, 152 milliliters of deionized water was added
to the alumina/ammonium-ferrierite mixture. Next, a mixture of 6.8 grams glacial acetic acid, 7.0 grams of citric acid and 152 milliliters of deionized water was slowly added to the alumina/ammonium-ferrierite mixture in the muller to peptize the alumina. The resulting alumina/ammonium-ferrierite/acid mixture was mulled for 10 minutes. Over a period of 15 minutes, a mixture of 0.20 grams of tetraamine palladium nitrate in 153 grams of deionized water was slowly added to mulled alumina/ammonium-ferrierite/acid mixture. The resulting mixture exhibited a 90:10 ratio of zeolite to alumina and a loss on ignition of 43.5%. The zeolite/alumina mixture was shaped by extruding the mixture through a stainless steel die plate (1/16” holes) of a 2.25 inch Bonnot extruder.

Six grams of the resulting zeolite/alumina mixture was impregnated with an aqueous solution of sodium hexachloroplatinate [IV] hexahydrate to incorporate 0.8 wt.% platinum into the 1/16 inch extrudate. The moist zeolite/alumina platinum impregnated extrudate was dried at 125°C for 2 hours in flowing air. The temperature was raised to a maximum temperature of 500°C and the zeolite/alumina platinum impregnated extrudate was calcined to yield a dehydrogenation-isomerization catalyst. The calcined catalyst was crushed and sized into 6-20 mesh particles before testing.

Dodecane was dehydrogenated and isomerized using the same reactor tube design as described in Example 3. A 16.1 weight percent conversion of dodecane to olefins was observed after twenty-fours hours at 459 °C. As tabulated in Table 7, of the olefins formed 86 weight percent were mono olefins, 1.2 weight percent were aromatic compounds and 6.8 weight percent were di-olefins. Of the total amount of olefins formed, 86 percent were branched, as determined by 1H NMR analysis.

**Table 7**

<table>
<thead>
<tr>
<th>Test Results</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Conversion (wt.% after 24 hours on-stream at 460 °C)</td>
<td>16.1</td>
</tr>
<tr>
<td>Temperature required for 16 wt. % conversion</td>
<td>459 °C</td>
</tr>
<tr>
<td>Selectivity to mono olefins at 16 wt. % conversion</td>
<td>86 wt.%</td>
</tr>
<tr>
<td>Selectivity to aromatics at 16 wt. % conversion</td>
<td>1.2 wt.%</td>
</tr>
<tr>
<td>Selectivity to di-olefins at 16 wt. % conversion</td>
<td>6.8 wt. %</td>
</tr>
<tr>
<td>% Branched C_{12} olefins in total C_{12} olefins</td>
<td>86</td>
</tr>
</tbody>
</table>
EXAMPLE 5: DEHYDROGENATION-ISOMERIZATION CATALYST. A zeolite portion of a dehydrogenation-isomerization catalyst was prepared as in Example 4. Six grams of the resulting zeolite/alumina mixture was impregnated with an aqueous solution of tetraamine palladium nitrate to incorporate 0.8 wt.% palladium into the 1/16 inch extrudates.

The moist zeolite/alumina palladium impregnated extrudate was dried at 125°C for 2 hours in flowing air. The temperature was raised to a maximum temperature of 500°C and the zeolite/alumina platinum impregnated extrudate was calcined to yield a dehydrogenation-isomerization catalyst. The calcined catalyst was carefully crushed and sized into 6-20 mesh particles before testing.

EXAMPLE 6: DEHYDROGENATION-ISOMERIZATION CATALYST: A dehydrogenation-isomerization catalyst was prepared according to the method for catalyst D of U.S. Patent No. 5,648,585 to Murray et al., entitled, “Process For Isomerizing Linear Olefins To Isoolefins.”

Ammonium-ferrierite having a molar silica to alumina ratio of 62:1, a surface area of 369 m2/g (P/Po=0.03), a soda content of 480 ppm wt and a n-hexane sorption capacity of 7.3 grams per 100 grams of zeolite was used. The catalyst components were mulled using a Lancaster mix muller. The mulled catalyst material was extruded using a Bonnot pin barrel extruder. The binder utilized was CATAPAL® D alumina from Sasol METHOCEL® F4M, hydroxypropyl methylcellulose, from The Dow Chemical Company was used as an extrusion aid.

The Lancaster mix muller was loaded with 632 grams of ammonium ferrierite (LOI of 3.4%) and 92 grams of CATAPAL®D alumina (LOI of 26.2%). The alumina was blended with the ferrierite for five minutes during which time 156 milliliters of de-ionized water was added. A mixture of 6.8 grams of glacial acetic acid and 156 milliliters of de-ionized water were added slowly to the muller in order to peptize the alumina. The mixture was mix-mulled for 10 minutes. Tetraamine platinum nitrate and Tetraamine palladium nitrate were added to the mix-muller in order to produce a catalyst that contained 0.25 wt. % palladium and 0.55 wt. % platinum. Ten grams of METHOCEL® F4M hydroxypropyl methylcellulose was added and the zeolite/alumina was mulled for 15 additional minutes. The extrudate was transferred to a Bonnot pin barrel extruder and extruded using a stainless
steel die plate with 1/16 inch holes. The extrudate was dried at 120 °C for 16 hours and then calcined in air at 500 °C for 2 hours. The calcined catalyst was carefully crushed and sized into 6-20 mesh particles before testing.
CLAIMS

1. A method for the production of aliphatic alcohols comprising:
   introducing a first hydrocarbon stream comprising olefins and paraffins into a
   hydrogenation unit, wherein the hydrogenation unit is configured to hydrogenate at least a
   portion of olefins in the first hydrocarbon stream to paraffins, and wherein at least a portion
   of the unreacted components of the first hydrocarbon stream and at least a portion of the
   hydrogenated olefins form a second hydrocarbon stream;
   introducing the second hydrocarbon stream into a dehydrogenation-isomerization
   unit, wherein the dehydrogenation-isomerization unit is configured to dehydrogenate at
   least a portion of the paraffins in the second hydrocarbon stream to olefins, and wherein the
   dehydrogenation-isomerization unit is further configured to isomerize at least a portion of
   linear olefins to branched olefins, and wherein at least a portion of the unreacted
   components of the second hydrocarbon stream and at least a portion of the products of the
   dehydrogenation and isomerization reactions form a third hydrocarbon stream, the third
   hydrocarbon stream comprising olefins and paraffins, and wherein at least a portion of the
   olefins in the third hydrocarbon stream is branched olefins; and,
   introducing at least a portion of the third hydrocarbon stream into a
   hydroformylation unit, wherein the hydroformylation unit is configured to hydroformylate
   at least a portion of the olefins in the third hydrocarbon stream to produce aliphatic
   alcohols, and wherein at least a portion of the produced aliphatic alcohols comprises a
   branched alkyl group.

2. The method of claim 1, wherein the first hydrocarbon stream is produced from a
   Fischer-Tropsch process.

3. The method of any one of claims 1 to 2, wherein the first hydrocarbon stream
   comprises olefins and paraffins having a carbon number of 7 to 18.

4. The method of any one of claims 1 to 3, wherein the hydrogenation unit is operated
   at a temperature range from about 175 °C to about 250 °C.

5. The method of any one of claims 1 to 4, wherein the hydrogenation unit is operated
   at a hydrogen flow rate between about 250 NL/L/hr and about 5000 NL/L/hr.
6. The method of any one of claims 1 to 5, wherein the hydrogenation unit is operated at a pressure range from about 10 atmospheres to about 50 atmospheres.

7. The method of any one of claims 1 to 6, wherein the dehydrogenation-isomerization unit is operated at a temperature range from about 300 °C to about 500 °C.

8. The method of any one of claims 1 to 7, wherein the dehydrogenation-isomerization unit is configured to operate at a pressure range from about 0.010 atmosphere to about 15 atmospheres.

9. The method of any one of claims 1 to 8, wherein a residence time of at least a portion of the second hydrocarbon stream in the dehydrogenation-isomerization unit is such that the conversion level of the paraffins in the second hydrocarbon stream composition to olefins is less than about 40 mole percent.

10. The method of any one of claims 1 to 9, wherein the hydroformylation unit is configured to produce greater than about 50 percent of aliphatic alcohols.

11. The method of any one of claims 1 to 10, wherein the hydroformylation unit is operated at a reaction temperature range from about 100 °C to about 300 °C.

12. The method of any one of claims 1 to 11, further comprising:
   forming a hydroformylation reaction stream, wherein the hydroformylation reaction stream comprises at least a portion of the unreacted components of the third hydrocarbon stream and at least a portion of the produced aliphatic alcohols;

   separating aliphatic alcohols from the hydroformylation reaction stream to produce a paraffins and unreacted olefins stream and an aliphatic alcohol product stream; and,

   introducing at least a portion of the paraffins and unreacted olefins stream into the dehydrogenation-isomerization unit.

13. The method of claim 12, wherein introducing at least a portion of the paraffins and unreacted olefins stream into the dehydrogenation-isomerization unit comprises combining at least a portion of the paraffins and unreacted olefins stream with at least a portion of the second hydrocarbon stream to produce a combined stream upstream of the dehydrogenation-isomerization unit and introducing at least a portion of the combined stream into the dehydrogenation-isomerization unit.
14. The method of any one of claims 1 to 13, further comprising:

introducing at least a portion of the produced aliphatic alcohols into a sulfation unit
and/or an oxyalkylation unit, wherein the sulfation unit is configured to sulfate at least a
portion of the aliphatic alcohols to produce aliphatic sulfates; wherein at least a portion of
the aliphatic sulfates produced comprises branched aliphatic sulfates and wherein the
oxyalkylation unit is configured to oxyalkylate at least a portion of the aliphatic alcohols to
produce oxyalkyl alcohols, wherein at least a portion of the oxyalkyl alcohols produced
comprises branched oxyalkyl alcohols.

15. The method of any one of claims 1 to 14, wherein the dehydrogenation-
isomerization unit comprises a dehydrogenation-isomerization catalyst configured to
catalyze both dehydrogenation reactions and isomerization reactions in the
dehydrogenation-isomerization unit, the catalyst comprising a hydrogen form of a zeolite
having a ferrierite isotypic framework structure, a binder, a coke-oxidizing compound, and
a paraffin dehydrogenation promoting compound.

16. The method of any one of claims 1 to 15, wherein the dehydrogenation-
isomerization unit comprises a plurality of zones, wherein the plurality of zones comprises
a first reaction zone and a second reaction zone, wherein the first reaction zone is
configured to dehydrogenate at least a portion of paraffins to olefins, and wherein the
second reaction zone is configured to isomerize at least a portion of linear olefins to
branched olefins, and wherein at least a portion of the unreacted components of the second
hydrocarbon stream and at least a portion of the products of the dehydrogenation and
isomerization reactions form a third hydrocarbon stream.

17. The method of claim 16, wherein the first reaction zone is operated at a temperature
range from about 300 °C to about 600 °C.

18. The method of claim 16, wherein the first reaction zone is operated at a total
reaction pressure between about 0.01 atmospheres and about 25.0 atmospheres.

19. The method of claim 16, wherein a residence time of at least a portion of the second
hydrocarbon stream in the first reaction zone is such that the conversion level of the
paraffins to olefins is less than about 50 mole percent.

20. The method of claim 16, wherein the second reaction zone is operated at a
temperature range from about 250 °C to about 500 °C.
21. The method of claim 16, wherein the second reaction zone is operated at a hydrocarbon partial pressure between about 0.1 atmosphere and about 10 atmospheres.

22. The method of any one of claims 16 to 21, further comprising introducing at least a portion of the second hydrocarbon stream exiting the first reaction zone into a heat exchanger, wherein the heat exchanger is configured to remove heat from a portion of the second hydrocarbon stream before it enters the second reaction zone.

23. The method of any one of claims 1 to 22, wherein the dehydrogenation-isomerization unit comprises a stacked bed catalyst configuration, wherein the stacked bed catalyst comprises a dehydrogenation catalyst and an isomerization catalyst.

24. A system for the production of aliphatic alcohols configured to perform the method according to any one of claims 1 to 23.
# INTERNATIONAL SEARCH REPORT

**International Application No**

PCT/US2004/034081

## A. CLASSIFICATION OF SUBJECT MATTER

<table>
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According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

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Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, CHEMABS Data, BEILSTEIN Data

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

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<td>US 4 438 287 A (IMAI ET AL)</td>
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**X** Further documents are listed in the continuation of box C

**X** Patent family members are listed in annex

* Special categories of cited documents

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-P document published prior to the international filing date but later than the priority date claimed

-T later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

-X document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

-V document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is considered in combination with one or more other documents, such combination being obvious to a person skilled in the art

-A document member of the same patent family

Date of the actual completion of the international search: 6 April 2005

Date of mailing of the international search report: 13/04/2005

Name and mailing address of the ISA

European Patent Office, P B 5818 Patentlaan 2 NL - 2280 HV Rijswijk

Tel: (+31-70) 340-2040, Tx: 31 551 epos nl, Fax: (+31-70) 340-3616

Authorized officer

Grammenoudi, S
<table>
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