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(54) SYSTEMS AND METHODS FOR PRODUCING A CRUDE PRODUCT

SYSTEME UND VERFAHREN ZUR HERSTELLUNG EINES ROHERZEUGNISSES

SYSTÈMES ET PROCÉDÉS DE PRODUCTION D'UN PRODUIT BRUT

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Description

TECHNICAL FIELD

5 **[0001]** The invention relates to systems and methods for treating or upgrading heavy oil feeds, and crude products produced using such systems and methods.

BACKGROUND

10 **[0002]** The petroleum industry is increasingly turning to heavy oil feeds such as heavy crudes, resids, coals, tar sands, etc. as sources for feedstocks. These feedstocks are characterized by high concentrations of asphaltenes rich residues, and low API gravities, with some being as low as less than 0° API.

[0003] PCT Patent Publication No. WO2008/014947, US Patent Publication No. 2008/0083650, US Patent Publication No. 2005/0241993, US Patent Publication No. 2007/0138057, and US Patent No. 6,660,157 describe processes, systems, and catalysts for processing heavy oil feeds. Heavy oil feedstock typically contains large levels of heavy metals. Some of the heavy metals such as nickel and vanadium tend to react quickly, leading to deposition or trapping of vanadium-rich solids in equipment such as reactors. The solid deposition reduces available volume for reaction, cutting down on run time.

[0004] US 2007/138056 A1 discloses a residuum full hydro-conversion slurry reactor system that allows the catalyst, unconverted oil, hydrogen and converted oil to circulate in a continuous mixture through the entire reactor with no confinement of the mixture.

[0005] There is still a need for improved systems and methods to upgrade / treat process heavy oil feeds with reduced build-ups of heavy metals in process equipment.

25 SUMMARY OF THE INVENTION

[0006] The essential technical features of the invention are explicitly defined in the wording of independent process claim 1 on file.

[0007] In one aspect, this invention relates to a process for by which a heavy oil feedstock can be upgraded. The process employs a plurality of contacting zones, separation zones and at least an interstage solvent deasphalting unit (SDA). The process comprises: a) combining a hydrogen containing gas feed, a heavy oil feedstock, and a slurry catalyst in a first contacting zone under hydrocracking conditions to convert at least a portion of the heavy oil feedstock to upgraded products; c) sending a mixture of the upgraded products, the slurry catalyst, the hydrogen containing gas, and unconverted heavy oil feedstock to a separation zone; d) in the separation zone, removing the upgraded products with the hydrogen containing gas as an overhead stream, and removing the slurry catalyst and the unconverted heavy oil feedstock as a non-volatile stream; e) sending at least a portion of the non-volatile stream to the SDA unit to separate the asphaltenes and slurry catalyst from the deasphalted oil; f) sending the deasphalted oil and the rest of the non-volatile stream from the previous separation zone to another contacting zone, under hydrocracking conditions with additional hydrogen gas and additional slurry catalyst, to convert the deasphalted oil to upgraded products; f) sending the upgraded products, the slurry catalyst, hydrogen, and unconverted deasphalted oil to a separation zone, whereby the upgraded products are removed with hydrogen as an overhead stream and the slurry catalyst and the unconverted deasphalted oil are removed as a non-volatile stream; and g) recycling to at least one of the contacting zones at least a portion of the non-volatile stream containing the slurry catalyst and the unconverted deasphalted oil.

[0008] In another aspect, there is provided a process employing a plurality of contacting zones, separation zones and at least an interstage solvent deasphalting unit (SDA) in which a heavy oil feedstock can be upgraded, and wherein at least a portion of the non-volatile stream from at least a contacting zone is sent to the SDA unit to separate the asphaltenes from the deasphalted oil.

[0009] In one aspect, this invention relates to a process for by which a heavy oil feedstock can be upgraded with reduced heavy metal deposits in the front-end contacting zones. The process employ a plurality of contacting zones and separation zones, the process comprising: a) combining a hydrogen containing gas feed, a heavy oil feedstock, and a slurry catalyst in a first contacting zone under hydrocracking conditions to convert at least a portion of the heavy oil feedstock to upgraded products, wherein water and / or steam being injected into first contacting zone in an amount of 1 to 25 weight % on the weight of the heavy oil feedstock; b) sending a mixture of the upgraded products, the slurry catalyst, the hydrogen containing gas, and unconverted heavy oil feedstock to a separation zone; c) in the separation zone, removing the upgraded products with the hydrogen containing gas as an overhead stream, and removing the slurry catalyst and the unconverted heavy oil feedstock as a non-volatile stream; d) sending the non-volatile stream to another contacting zone under hydrocracking conditions with additional hydrogen gas, unconverted heavy oil feedstock, and optionally, a fresh slurry catalyst to convert the unconverted heavy oil feedstock to upgraded products; f) sending

the upgraded products, the slurry catalyst, hydrogen, and unconverted heavy oil feedstock to a separation zone, whereby the upgraded products are removed with hydrogen as an overhead stream and the slurry catalyst and the unconverted heavy oil feedstock are removed as a non-volatile stream; and g) recycling to at least one of the contacting zones at least a portion of the non-volatile stream.

[0010] In another aspect, the invention relates to a method for upgrading a heavy oil feedstock employing a plurality of contacting zones and separation zones, in which water and / or steam is injected into the first contacting zone, and wherein at least a portion of the non-volatile stream from a separation zone other than the first separation zone is recycled to the first contacting zone, wherein the recycled stream ranges between 35 to 50 wt. % of the total heavy oil feedstock to the process.

[0011] In one aspect, this invention relates to a process for by which a heavy oil feedstock can be upgraded. The process employs a plurality of contacting zones and separation zones, the process comprising: a) a heavy oil feedstock with at least a portion of the heavy oil feedstock is fed to a contacting zone other than the first contacting zone; b) combining a hydrogen containing gas feed, a portion of the heavy oil feedstock, and a slurry catalyst in a first contacting zone under hydrocracking conditions to convert at least a portion of the heavy oil feedstock to upgraded products; c) sending a mixture of the upgraded products, the slurry catalyst, the hydrogen containing gas, and unconverted heavy oil feedstock to a separation zone; d) in the separation zone, removing the upgraded products with the hydrogen containing gas as an overhead stream, and removing the slurry catalyst and the unconverted heavy oil feedstock as a non-volatile stream; e) sending the non-volatile stream to another contacting zone under hydrocracking conditions with additional hydrogen gas, at least a portion of the heavy oil feedstock, and optionally, fresh slurry catalyst to convert the unconverted heavy oil feedstock to upgraded products; f) sending the upgraded products, the slurry catalyst, hydrogen, and unconverted heavy oil feedstock to a separation zone, whereby the upgraded products are removed with hydrogen as an overhead stream and the slurry catalyst and the unconverted heavy oil feedstock are removed as a non-volatile stream; and g) recycling to the first contacting zone at least a portion of the non-volatile stream.

[0012] In another aspect, the process employs a plurality of contacting zones and separation zones, the process comprising: a) providing a slurry catalyst containing a used slurry catalyst and optionally a fresh catalyst slurry feed; b) combining a hydrogen containing gas feed, the heavy oil feedstock, and the slurry catalyst in a contacting zone under hydrocracking conditions to convert at least a portion of the heavy oil feedstock to upgraded products; c) sending a mixture comprising the upgraded products, the slurry catalyst, the hydrogen containing gas, and unconverted heavy oil feedstock to a separation zone; d) in the separation zone, removing the upgraded products with the hydrogen containing gas as an overhead stream, and removing the slurry catalyst and the unconverted heavy oil feedstock as a non-volatile stream; e) sending the non-volatile stream to another contacting zone under hydrocracking conditions with additional hydrogen gas and a fresh slurry catalyst to convert the unconverted heavy oil feedstock to upgraded products; f) sending the upgraded products, the slurry catalyst, hydrogen, and unconverted heavy oil feedstock to a separation zone, whereby the upgraded products are removed with hydrogen as an overhead stream and the slurry catalyst and the unconverted heavy oil feedstock are removed as a non-volatile stream; and g) recycling to the first contacting zone at least a portion of the non-volatile stream.

[0013] In yet another aspect, there is provided a process employing a plurality of contacting zones and separation zones in which a heavy oil feedstock can be upgraded, and wherein the fresh slurry catalyst is split between the contacting zones.

[0014] In one aspect, the process employs a plurality of contacting zones and separation zones, the process comprising: a) combining a hydrogen containing gas feed, a heavy oil feedstock, and a slurry catalyst in a first contacting zone under hydrocracking conditions to convert at least a portion of the heavy oil feedstock to upgraded products; b) sending a mixture of the upgraded products, the slurry catalyst, the hydrogen containing gas, and unconverted heavy oil feedstock to a separation zone; c) in the separation zone, removing the upgraded products with the hydrogen containing gas as an overhead stream, and removing the slurry catalyst and the unconverted heavy oil feedstock as a non-volatile stream; d) sending the non-volatile stream to another contacting zone under hydrocracking conditions with additional hydrogen gas, unconverted heavy oil feedstock, and optionally, a fresh slurry catalyst to convert the unconverted heavy oil feedstock to upgraded products; f) sending the upgraded products, the slurry catalyst, hydrogen, and unconverted heavy oil feedstock to a separation zone, whereby the upgraded products are removed with hydrogen as an overhead stream and the slurry catalyst and the unconverted heavy oil feedstock are removed as a non-volatile stream; and wherein the first contacting zone operates at a temperature of at least at least 10°F (5.65°C) lower than the next contacting zone in series.

[0015] In yet another aspect, the invention relates to a process for by which a heavy oil feedstock can be upgraded with reduced heavy metal deposits in the front-end contacting zones. The process employ a plurality of contacting zones and separation zones, comprising: a) combining a hydrogen containing gas feed, a heavy oil feedstock, and a slurry catalyst in a first contacting zone under hydrocracking conditions to convert at least a portion of the heavy oil feedstock to upgraded products; b) sending a mixture of the upgraded products, the slurry catalyst, the hydrogen containing gas, and unconverted heavy oil feedstock to a separation zone; c) in the separation zone, removing the upgraded products

with the hydrogen containing gas as an overhead stream, and removing the slurry catalyst and the unconverted heavy oil feedstock as a non-volatile stream; d) sending the non-volatile stream to another contacting zone under hydrocracking conditions with additional hydrogen gas, unconverted heavy oil feedstock, and optionally, a fresh slurry catalyst to convert the unconverted heavy oil feedstock to upgraded products; f) sending the upgraded products, the slurry catalyst, hydrogen, and unconverted heavy oil feedstock to a separation zone, wherein the upgraded products are removed with hydrogen as an overhead stream and the slurry catalyst and the unconverted heavy oil feedstock are removed as a non-volatile stream; and wherein the slurry catalyst to the first separation zone comprises at least a portion of a non-volatile stream from one of the separation zones as a recycled catalyst stream, and wherein the recycled catalyst stream is between 35 to 50 wt. % of the heavy oil feedstock

BRIEF DESCRIPTION OF THE DRAWINGS

[0016]

Figure 1 is a block diagram that schematically illustrates an embodiment of a hydroprocessing system for upgrading a heavy oil feedstock, with a plurality of contacting zones and separation zones, wherein water and or steam is injected into the front end contacting zone.

Figure 2 is a flow diagram of a process to upgrade heavy oil feeds with water injection.

Figure 3 is a flow diagram of a process to upgrade heavy oil feeds with steam injection directly into a front end contacting zone.

Figure 4 is a flow diagram of another embodiment of process to upgrade heavy oil feeds with a recycled catalyst stream at a sufficient rate to reduce heavy metal build-up.

Figure 5 is a block diagram that schematically illustrates an embodiment of a hydroprocessing system for upgrading a heavy oil feedstock, having a split fresh catalyst feed scheme, a split heavy oil feed scheme, and additional interstage hydrocarbon oil feedstock.

Figure 6 is a block diagram that schematically illustrates another embodiment of a hydroprocessing system for upgrading a heavy oil feedstock with a solvent deasphalting unit for pre-treating the heavy oil feedstock.

Figure 7 is a flow diagram of a process to upgrade heavy oil feeds with an embodiment of the catalyst split feed scheme, wherein fresh catalyst feed is fed into all reactors in the process.

Figure 8 is a flow diagram of a process to upgrade heavy oil feeds wherein the fresh catalyst feed is diverted from the first reactor to other reactors in the process, and wherein optional / additional hydrocarbon oil is fed to the reactors as feedstock.

Figure 9 is a flow diagram of another embodiment of a process to upgrade heavy oil feeds, wherein all of the fresh catalyst feed is sent to the last reactor in the process.

Figure 10 is a flow diagram of another embodiment of a process to upgrade heavy oil feeds, wherein some of the untreated heavy oil feed is diverted from the first reactor sent to other reactors in the process.

DETAILED DESCRIPTION

[0017] The following terms will be used throughout the specification and will have the following meanings unless otherwise indicated.

[0018] As used herein, "heavy oil" feed or feedstock refers to heavy and ultra-heavy crudes, including but not limited to resids, coals, bitumen, shale oils, tar sands, etc. Heavy oil feedstock may be liquid, semi-solid, and / or solid. Examples of heavy oil feedstock that might be upgraded as described herein include but are not limited to Canada Tar sands, vacuum resid from Brazilian Santos and Campos basins, Egyptian Gulf of Suez, Chad, Venezuelan Zulia, Malaysia, and Indonesia Sumatra. Other examples of heavy oil feedstock include bottom of the barrel and residuum left over from refinery processes, including "bottom of the barrel" and "residuum" (or "resid") -- atmospheric tower bottoms, which have a boiling point of at least 343°C. (650°F.), or vacuum tower bottoms, which have a boiling point of at least 524°C. (975°F.), or "resid pitch" and "vacuum residue" - which have a boiling point of 524°C. (975°F.) or greater.

[0019] Properties of heavy oil feedstock may include, but are not limited to: TAN of at least 0.1, at least 0.3, or at least 1; viscosity of at least 10 cSt; API gravity at most 15 in one embodiment, and at most 10 in another embodiment. A gram of heavy oil feedstock typically contains at least 0.0001 grams of Ni/V/Fe; at least 0.005 grams of heteroatoms; at least 0.01 grams of residue; at least 0.04 grams C5 asphaltenes; at least 0.002 grams of MCR; per gram of crude; at least 0.00001 grams of alkali metal salts of one or more organic acids; and at least 0.005 grams of sulfur. In one embodiment, the heavy oil feedstock has a sulfur content of at least 5 wt. % and an API gravity of from -6 to +6.

[0020] The terms "treatment," "treated," "upgrade", "upgrading" and "upgraded", when used in conjunction with a heavy oil feedstock, describes a heavy oil feedstock that is being or has been subjected to hydroprocessing, or a resulting material or crude product, having a reduction in the molecular weight of the heavy oil feedstock, a reduction in the boiling

point range of the heavy oil feedstock, a reduction in the concentration of asphaltenes, a reduction in the concentration of hydrocarbon free radicals, and/or a reduction in the quantity of impurities, such as sulfur, nitrogen, oxygen, halides, and metals.

[0021] The upgrade or treatment of heavy oil feeds is generally referred herein as "hydroprocessing". Hydroprocessing is meant as any process that is carried out in the presence of hydrogen, including, but not limited to, hydroconversion, hydrocracking, hydrogenation, hydrotreating, hydrodesulfurization, hydrodenitrogenation, hydrodemetallation, hydrodearomatization, hydroisomerization, hydrodewaxing and hydrocracking including selective hydrocracking. The products of hydroprocessing may show improved viscosities, viscosity indices, saturates content, low temperature properties, volatilities and depolarization, etc.

[0022] As used herein, hydrogen refers to hydrogen, and / or a compound or compounds that when in the presence of a heavy oil feed and a catalyst react to provide hydrogen.

[0023] SCF / BBL (or scf / bbl) refers to a unit of standard cubic foot of gas (N₂, H₂, etc.) per barrel of hydrocarbon feed.

[0024] Nm³/m³ refers to normal cubic meters of gas per cubic meter of heavy oil feed.

[0025] VGO or vacuum gas oil, referring to hydrocarbons with a boiling range distribution between 343°C (650°F) and 538°C (1000°F) at 0.101 MPa.

[0026] "wppm" means weight parts per million.

[0027] As used herein, the term "catalyst precursor" refers to a compound containing one or more catalytically active metals, from which compound a catalyst is eventually formed. It should be noted that a catalyst precursor may be catalytically active as a hydroprocessing catalyst. As used herein, "catalyst precursor" may be referred herein as "catalyst" when used in the context of a catalyst feed.

[0028] As used herein, the term "used catalyst" refers to a catalyst that has been used in at least a reactor in a hydroprocessing operation and whose activity has thereby been diminished. For example, if a reaction rate constant of a fresh catalyst at a specific temperature is assumed to be 100%, the reaction rate constant for a used catalyst is 95% or less in one embodiment, 80% or less in another embodiment, and 70% or less in a third embodiment. The term "used catalyst" may be used interchangeably with "recycled catalyst," "used slurry catalyst" or "recycled slurry catalyst."

[0029] As used herein, the term "fresh catalyst" refers to a catalyst or a catalyst precursor that has not been used in a reactor in a hydroprocessing operation. The term fresh catalyst herein also includes "re-generated" or "rehabilitated" catalysts, i.e., catalyst that has been used in at least a reactor in a hydroprocessing operation ("used catalyst") but its catalytic activity has been restored or at least increased to a level well above the used catalytic activity level. The term "fresh catalyst" may be used interchangeably with "fresh slurry catalyst."

[0030] As used herein, the term "slurry catalyst" (or sometimes referred to as "slurry", or "dispersed catalyst") refers to a liquid medium, e.g., oil, water, or mixtures thereof, in which catalyst and / or catalyst precursor particles (particulates or crystallites) having very small average dimensions are dispersed within.

[0031] As used herein, the "catalyst feed" includes any catalyst suitable for upgrading heavy oil feed stocks, e.g., one or more bulk catalysts and / or one or more catalysts on a support. The catalyst feed may include at least a fresh catalyst, a used catalyst only, or mixtures of at least a fresh catalyst and a used catalyst. In one embodiment, the catalyst feed is in the form of a slurry catalyst.

[0032] As used herein, the term "bulk catalyst" may be used interchangeably with "unsupported catalyst," meaning that the catalyst composition is NOT of the conventional catalyst form which has, i.e., having a preformed, shaped catalyst support which is then loaded with metals via impregnation or deposition catalyst. In one embodiment, the bulk catalyst is formed through precipitation. In another embodiment, the bulk catalyst has a binder incorporated into the catalyst composition. In yet another embodiment, the bulk catalyst is formed from metal compounds and without any binder. In a fourth embodiment, the bulk catalyst is a dispersing-type catalyst for use as dispersed catalyst particles in mixture of liquid (e.g., hydrocarbon oil). In one embodiment, the catalyst comprises one or more commercially known catalysts, e.g., Microcat™ from ExxonMobil Corp.

[0033] As used herein, the term "contacting zone" refers to an equipment in which the heavy oil feed is treated or upgraded by contact with a slurry catalyst feed in the presence of hydrogen. In a contacting zone, at least a property of the crude feed may be changed or upgraded. The contacting zone can be a reactor, a portion of a reactor, multiple portions of a reactor, or combinations thereof. The term "contacting zone" may be used interchangeably with "reacting zone."

[0034] As used herein, the term "separation zone" refers to an equipment in which upgraded heavy oil feed from a contacting zone is either fed directly into, or subjected to one or more intermediate processes and then fed directly into the separation zone, e.g., a flash drum or a high pressure separator, wherein gases and volatile liquids are separated from the non-volatile fraction. In one embodiment, the non-volatile fraction stream comprises unconverted heavy oil feed, a small amount of heavier hydrocracked liquid products (synthetic or less-volatile / non-volatile upgraded products), the slurry catalyst and any entrained solids (asphaltenes, coke, etc.).

[0035] As used herein, the term "bleed stream" or "bleed off stream" refers to a stream containing used (or recycled) catalyst, being "bled" or diverted from the hydroprocessing system, helping to prevent or "flush" accumulating metallic

sulfides and other unwanted impurities from the upgrade system.

[0036] The present invention relates to an improved system to treat or upgrade heavy oil feeds, particularly heavy oil feedstock having high levels of heavy metals.

[0037] In a typical prior art hydroprocessing system having a plurality of contacting zones (reactors) in series, it is observed that the feed stream to the 2nd contacting zone should generally be cleaner than heavy oil feed into the first contacting zone in the system, i.e., having less impurities such as nickel, vanadium, nitrogen, sulfur, etc., as the heavy oil has gone through a treatment process in the first contacting zone. It is also observed that the feed stream into the last contacting zone in the system should generally be cleaner than the feed stream to the prior contacting zone(s) in the system.

[0038] In a typical hydroprocessing system, it has been further observed that in the catalyst feed scheme of the prior art, the feed streams to the subsequent contacting zones in the system are typically more concentrated in terms of certain impurities, e.g., MCR, C₅ and C₇ asphaltenes contents, etc., thus promoting coke formation in the latter contacting zones in the system.

[0039] It has also been observed that the feed stream to subsequent contacting zones in the system has properties different than the properties of the heavy oil feed to the preceding contacting zone(s) in the system, including: a) lower TAN; b) viscosity; c) lower residue content; d) lower API gravity; e) lower content of metals in metal salts of organic acids; and g) combinations thereof. However, it has also been observed that it is generally more difficult to process the feed to the subsequent contacting zones in the system in terms of the conversion rate and / or the properties of the resulting crude product. Additionally with the prior art feeding scheme (fresh catalyst going to the 1st contacting zone), it is observed that there is more coke formation in the subsequent contacting zones than in the 1st contacting zone. It is speculated that the coke formation perhaps has something to do with the more-difficult-to-process feed to the subsequent contacting zones and / or the reduced activity of the catalyst feed to the subsequent contacting zones.

[0040] In one embodiment, the upgrade process comprises a plurality of reactors for contacting zones, with the reactors being the same or different in configurations. Examples of reactors that can be used herein include stacked bed reactors, fixed bed reactors, ebullating bed reactors, continuous stirred tank reactors, fluidized bed reactors, spray reactors, liquid / liquid contactors, slurry reactors, liquid recirculation reactors, and combinations thereof. In one embodiment, the reactor is an up-flow reactor. In another embodiment, a down-flow reactor. In one embodiment, the contacting zone refers to at least a slurry-bed hydrocracking reactor in series with at least a fixed bed hydrotreating reactor. In another embodiment, at least one of the contacting zones further comprises an in-line hydrotreater, capable of removing removed over 70% of the sulfur, over 90% of nitrogen, and over 90% of the heteroatoms in the crude product being processed.

[0041] In one embodiment, the contacting zone comprises a plurality of reactors in series, providing a total residence time ranging from 0.1 to 15 hours. In a second embodiment, the resident time ranges from 0.5 to 5 hrs. In a third embodiment, the total residence time in the contacting zone ranges from 0.2 to 2 hours.

[0042] Depending on the conditions and location of the separation zone, in one embodiment, the amount of heavier hydrocracked products in the non-volatile fraction stream is less than 50 wt. % (of the total weight of the non-volatile stream). In a second embodiment, the amount of heavier hydrocracked products in the non-volatile stream from the separation zone is less than 25 wt. %. In a third embodiment, the amount of heavier hydrocracked products in the non-volatile stream from the separation zone is less than 15 wt. %. It should be noted that at least a portion of the slurry catalyst remains with the upgraded feedstock as the upgraded materials is withdrawn from the contacting zone and fed into the separation zone, and the slurry catalyst continues to be available in the separation zone and exits the separation zone with the non-volatile liquid fraction.

[0043] In one embodiment, both the contacting zone and the separation zone are combined into one equipment, e.g., a reactor having an internal separator, or a multi-stage reactor-separator. In this type of reactor-separator configuration, the vapor product exits the top of the equipment, and the non-volatile fractions exit the side or bottom of the equipment with the slurry catalyst and entrained solid fraction, if any.

[0044] In one embodiment, the slurry catalyst stream contains a fresh catalyst. In another embodiment, the slurry catalyst stream contains a mixture of at least a fresh catalyst and a recycled (used) catalyst. In a third embodiment, the slurry catalyst stream comprises a used catalyst. In another embodiment, the slurry catalyst contains a well-dispersed catalyst precursor composition capable of forming an active catalyst in situ within the feed heaters and/or the contacting zone. The catalyst particles can be introduced into the medium (diluent) as powder in one embodiment, a precursor in another embodiment, or after a pretreatment step in a third embodiment. In one embodiment, the medium (or diluent) is a hydrocarbon oil diluent. In another embodiment, the liquid medium is the heavy oil feedstock itself. In yet another embodiment, the liquid medium is a hydrocarbon oil other than the heavy oil feedstock, e.g., a VGO medium or diluent.

[0045] In one embodiment, the bleed off stream comprises non-volatile materials from a separation zone in the system, typically the last separation zone, comprising unconverted materials, slurry catalyst, a small amount of heavier hydrocracked liquid products, small amounts of coke, asphaltenes, etc. In another embodiment, the bleed off stream is the bottom stream from an interstage solvent deasphalting unit in the system. In embodiments wherein the bleed off stream is diverted from the bottom stream of a separation zone, the bleed stream typically ranges from 1 to 35 wt. %; 3-20 wt.

%; or 5-15wt. % of the total heavy oil feedstock to the system. In embodiments therein the bleed off stream is diverted from the bottom of a deasphalting unit, the bleed off stream ranges from 0.30 to 5 wt.%; 1-30 wt. %; or 0.5 to 10 wt. % of the heavy oil feed stock.

[0046] In one embodiment, the bleed-off stream contains between 3 to 30 wt. % slurry catalyst. In another embodiment, the slurry catalyst amount ranges from 5 to 20 wt. %. In yet another embodiment, the bleed-off stream contains an amount of slurry catalyst ranging from 1 to 15 wt. % in concentration.

[0047] In some embodiments, instead of sending all of the fresh catalyst to the first contacting zone as in the prior art process, at least a portion of the fresh catalyst is diverted to at least one other contacting zones (other than the 1st contacting zone) in the system.

[0048] Also in some embodiments, instead of sending all of the heavy oil feed to be upgraded to the first contacting zone, at least a portion of the heavy oil feed is diverted to at least one other contacting zones in the system.

[0049] In other embodiments, a combination feed scheme is employed with a portion of the fresh catalyst feed and a portion of the heavy oil feed being diverted to at least one other contact zones other than the first contacting zone in the heavy oil upgrading system.

[0050] In one embodiment, the upgrade system comprises at least two upflow reactors in series with at least two separators, with each separator being positioned right after each reactor and with the interstage SDA unit being positioned before at least one reactor in the system. In another embodiment, the upgrade system comprises at least two upflow reactors and at least two separators in series, with of each of the separators being positioned right after each reactor, and the interstage SDA unit being positioned after the 1st separator in the series. In a fourth embodiment, the upgrade system may comprise a combination of separate reactors and separate separators in series with multi-stage reactor-separators, with the SDA being positioned as an interstage treatment system between any two reactors in series.

[0051] Heavy Oil Feed: The unconverted heavy oil feed here herein may comprise one or more different heavy oil feeds from different sources as a single feed stream, or as separate heavy oil feed streams. In some embodiments of the present invention, at least a portion of the heavy oil feed (to be upgraded) is "split" or diverted to at least one other contacting zones in the system (other than the first contacting zone), or to the interstage SDA unit prior to being fed into a contacting zone.

[0052] In one embodiment, "at least a portion" means at least 5% of the heavy oil feed to be upgraded is diverted to at least one other contacting zones in the system other than the first contacting zone. In another embodiment, at least 10%. In a third embodiment, at least 20%. In a fourth embodiment, at least 30% of the heavy oil feed is diverted to at least a contacting zone other than the first one in the system. In one embodiment, the heavy oil feedstock is preheated prior to being blended with the slurry catalyst feed stream(s). In another embodiment, the blend of heavy oil feedstock and slurry catalyst feed is preheated to create a feedstock that is sufficiently of low viscosity to allow good mixing of the catalyst into the feedstock. In one embodiment, the preheating is conducted at a temperature that is at least about 100°C (180°F) less than the hydrocracking temperature within the contacting zone. In another embodiment, the preheating is at a temperature that is about at least 50°C less than the hydrocracking temperature within the contacting zone.

[0053] Additional Hydrocarbon Feed: In one embodiment, additional hydrocarbon oil feed, e.g., VGO (vacuum gas oil), naphtha, MCO (medium cycle oil), solvent donor, or other aromatic solvents, etc. in an amount ranging from 2 to 40 wt. % of the heavy oil feed can be optionally added as part of the heavy oil feed stream to any of the contacting zones in the system. In one embodiment, the additional hydrocarbon feed functions as a diluent to lower the viscosity of the heavy oil feed.

[0054] Embodiments of The Heavy Oil Split Feed Scheme: In some embodiments, at least a portion of the heavy oil feed (to be upgraded) is "split" or diverted to at least one other contacting zones in the system (other than the first contacting zone).

[0055] In one embodiment, "at least a portion" meaning at least 5% of the heavy oil feed to be upgraded. In another embodiment, at least 10%. In a third embodiment, at least 20%. In a fourth embodiment, at least 30% of the heavy oil feed is diverted to at least a contacting zone other than the first one in the system.

[0056] In one embodiment, less than 90% of the unconverted heavy oil feed is fed to the first reactor in the system, with 10% or more of the unconverted heavy oil feed being diverted to the other contacting zone(s) in the system. In another embodiment, the heavy oil feed is being equally split between the contacting zones in the system. In yet another embodiment, less than 80% of the unconverted heavy oil feed is fed to the first contacting zone in the system, and the remaining heavy oil feed is diverted to the last contacting zone in the system. In a fourth embodiment, less than 60% of the heavy oil feed is fed to the first contacting zone in the system, and the remainder of the unconverted heavy oil feed is equally split between the other contacting zones in the system.

[0057] The unconverted heavy oil feed herein may comprise one or more different heavy oil feeds from different sources as a single feed stream or separate heavy oil feed streams. In one embodiment, a single heavy oil conduit pipe goes to all the contacting zones. In another embodiment, multiple heavy oil conduits are employed to supply the heavy oil feed to the different contacting zones, with some heavy oil feed stream(s) going to one or more contacting zones, and some of the other unconverted heavy oil feed stream(s) going to one or more different contacting zones.

[0058] In one embodiment, the heavy oil feedstock is preheated prior to being blended with the slurry catalyst feed, and / or prior to being introduced into the hydrocracking reactors (contacting zones). In another embodiment, the blend of heavy oil feedstock and slurry catalyst feed is preheated to create a feedstock that is sufficiently of low viscosity to allow good mixing of the catalyst into the feedstock.

[0059] In one embodiment, the preheating is conducted at a temperature that is about 100°C (180°F) less than the hydrocracking temperature within the contacting zone. In another embodiment, the preheating is at a temperature that is about 50°C less than the hydrocracking temperature within the contacting zone.

[0060] Hydrogen Feed: In one embodiment, a hydrogen containing gas is provided to the process. The hydrogen can also be added to the heavy oil feed prior to entering the preheater, or after the preheater. In one embodiment, the hydrogen feed enters the contacting zone co-currently with the heavy oil feed in the same conduit. In another embodiment, the hydrogen source may be added to the contacting zone in a direction that is counter to the flow of the crude feed. In a third embodiment, the hydrogen enters the contacting zone via a gas conduit separately from the combined heavy oil and slurry catalyst feed stream. In a fourth embodiment, the hydrogen feed is introduced directly to the combined catalyst and heavy oil feedstock prior to being introduced into the contacting zone. In yet another embodiment, the hydrogen gas and the combined heavy oil and catalyst feed are introduced at the bottom of the reactor as separate streams. In yet another embodiment, hydrogen gas can be fed to several sections of the contacting zone.

[0061] In one embodiment, the hydrogen source is provided to the process at a rate (based on ratio of the gaseous hydrogen source to the crude feed) of 0.1 Nm³/m³ to about 100,000 Nm³/m³ (0.563 to 563,380 SCF/bbl), about 0.5 Nm³/m³ to about 10,000 Nm³/m³ (2.82 to 56,338 SCF/bbl), about 1 Nm³/m³ to about 8,000 Nm³/m³ (5.63 to 45,070 SCF/bbl), about 2 Nm³/m³ to about 5,000 Nm³/m³ (11.27 to 28,169 SCF/bbl), about 5 Nm³/m³ to about 3,000 Nm³/m³ (28.2 to 16,901 SCF/bbl), or about 10 Nm³/m³ to about 800 Nm³/m³ (56.3 to 4,507 SCF/bbl). In one embodiment, some of the hydrogen (25 -75%) is supplied to the first contacting zone, and the rest is added as supplemental hydrogen to other contacting zones in system.

[0062] In one embodiment, the upgrade system produces a volume yield over 100% (compared to the heavy oil input) in upgraded products as added hydrogen expands the heavy oil total volume. The upgraded products, i.e., lower boiling hydrocarbons, in one embodiment include liquefied petroleum gas (LPG), gasoline, diesel, vacuum gas oil (VGO), and jet and fuel oils. In a second embodiment, the upgrade system provides a volume yield of at least 110% in the form of LPG, naphtha, jet & fuel oils, and VGO. In a third embodiment, over 115%.

[0063] In one embodiment of the upgrade system, at least 98 wt % of heavy oil feed is converted to lighter products. In a second embodiment, at least 98.5% of heavy oil feed is converted to lighter products. In a third embodiment, the conversion rate is at least 99%. In a fourth embodiment, the conversion rate is at least 95%. In a fifth embodiment, the conversion rate is at least 80%. In a sixth embodiment, the conversion rate is at least 60%. As used herein, conversion rate refers to the conversion of heavy oil feedstock to less than 1000°F. (538°C) boiling point materials.

[0064] The hydrogen source, in some embodiments, is combined with carrier gas(es) and recirculated through the contacting zone. Carrier gas may be, for example, nitrogen, helium, and/or argon. The carrier gas may facilitate flow of the crude feed and/or flow of the hydrogen source in the contacting zone(s). The carrier gas may also enhance mixing in the contacting zone(s). In some embodiments, a hydrogen source (for example, hydrogen, methane or ethane) may be used as a carrier gas and recirculated through the contacting zone.

[0065] Catalyst Feed: In one embodiment, all of the slurry catalyst feed is provided to the first contacting zone. In other embodiments, at least a portion of the catalyst feed is "split" or diverted to at least one other contacting zones in the system (other than the first contacting zone). In another embodiment, all the contacting zones in operation receive a slurry catalyst feed (along with a heavy oil feed). In yet another embodiment, the process is configured for a flexible catalyst feed scheme such that the fresh catalyst can sometimes be fed entirely to the last reactor in the system for certain process conditions (for certain desired product characteristics), or 50% to the first reactor in the system for some of the process runs, or split equally or according to pre-determined proportions to all of the reactors in the system, or split according to pre-determined proportions for the same fresh catalyst to be fed to the different reactors at different concentrations.

[0066] The slurry catalyst feed used herein may comprise one or more different slurry catalysts as a single catalyst feed stream or separate feed streams. In one embodiment, a single fresh catalyst feed stream is supplied to the contacting zones. In another embodiment, the catalyst feed comprises multiple and different catalyst types, with a certain catalyst type going to one or more contacting zones (e.g., the first contacting zone in the system) as a separate stream, and a different slurry catalyst going to contacting zone(s) other than the 1st contacting zone in the system as a different catalyst stream.

[0067] In one embodiment, "at least a portion" means at least 10% of the fresh catalyst. In another embodiment, at least 20%. In a third embodiment, at least 40%. In a fourth embodiment, at least 50% of the fresh catalyst is diverted to at least a contacting zone other than the first one in the system. In a fifth embodiment, all of the fresh catalyst is diverted to a contacting zone other than the 1st contacting zone.

[0068] In one embodiment, less than 20% of the fresh catalyst is fed to the first reactor in the system, with 80% or

more of the fresh catalyst being diverted to the other contacting zone(s) in the system. In another embodiment, the fresh catalyst is being equally split between the contacting zones in the system. In one embodiment, at least a portion of the fresh catalyst feed is sent to at least one of the intermediate contacting zones and / or the last contacting zone in the system. In another embodiment, all of the fresh catalyst is sent to the last contacting zone in the system, with the first contacting zone in the system only getting recycled catalyst from one or more of the processes in the system, e.g., from one of the separation zones in the system or from a solvent deasphalting unit.

[0069] In one embodiment with an interstage SDA unit, at least a portion of the fresh catalyst feed is sent to the contacting zone immediately following the interstage SDA unit. In another embodiment, all of the fresh catalyst is sent to contacting zone(s) other than the 1st one in the system, with the first contacting zone only getting SDA bottoms from the SDA unit and recycled catalyst from one or more of the processes in the system, e.g., from one of the separation zones in the system.

[0070] In one embodiment, the fresh catalyst is combined with the recycled catalyst stream from one of the processes in the system, e.g., a separation zone, a distillation column, a SDA unit, or a flash tank, and the combined catalyst feed is thereafter blended with heavy oil feedstock for feeding into the contacting zone(s). In another embodiment, the fresh catalyst and the recycled catalyst streams are blended into the heavy oil feedstock as separate streams.

[0071] In one embodiment, the recycled catalyst stream from one of the processes in the system, e.g., a separation zone, the SDA unit, etc., is combined with fresh slurry catalyst as one single catalyst feed stream. The combined catalyst feed is thereafter blended with the (treated or untreated) heavy oil feedstock stream(s) for feeding into the contacting zone(s). In another embodiment, the fresh catalyst and the recycled catalyst streams are blended into the heavy oil feedstock stream(s) as separate streams.

[0072] In one embodiment, the process is configured for a flexible catalyst feed scheme such that the catalyst feed can sometimes be fed at full rate (100% of the required catalyst rate) to the first reactor in the system for a certain period of time, then split equally or according to pre-determined proportions to all of the reactors in the system for a pre-determined amount of time, or split according to pre-determined proportions for the catalyst feed to be fed to the different reactors at different concentrations.

[0073] In one embodiment, sending different catalysts to the front end and back end contacting zones can be useful in mitigating the vanadium trapping issue and sustain the overall upgrade performance. In one embodiment, a Ni-only or a NiMo sulfide slurry catalyst rich in Ni is sent to the front end reactor to help reduce vanadium trapping in the system, while a different catalyst, e.g., Mo sulfide or a NiMo sulfide catalyst rich in Mo, can be injected into the back end reactor(s) to maintain an overall high conversion rate, improve product quality and possibly reduce the gas yield in one embodiment. As used herein, a slurry catalyst rich in Ni means that the Ni / Mo ratio is greater than 0.15 (as wt. %) Conversely, a slurry catalyst rich in Mo means that the Ni / Mo ratio is less than 0.05 (as wt. %).

[0074] In one embodiment, the slurry catalyst feed is first preconditioned before entering one of the contacting zones, or before being brought into contact with the heavy oil feed before entering the contacting zones. In one example, the catalyst enters into a preconditioning unit along with hydrogen at a rate from 89 l./l. to 1336 l./l. (500 to 7500 SCF/BBL) wherein BBL here refers to the total volume of heavy oil feed to the system. It is believed that instead of bringing a cold catalyst in contact with the heavy oil feed, the preconditioning step helps with the hydrogen adsorption into the active catalyst sites, and ultimately the conversion rate. In one embodiment in the precondition unit, the slurry catalyst / hydrogen mixture is heated to a temperature between 300°F to 1000°F (149 to 538°C). In another embodiment, the catalyst is preconditioned in hydrogen at a temperature of 500 to 725 °F (260 to 385°C.). In yet another embodiment, the mixture is heated under a pressure of 2068.4 to 2206.2 kPa (300 to 3200 psi) in one embodiment; 3447.4 to 20684.3 kPa (500 to 3000 psi) in a second embodiment; and 4136.8 to 17246.9 kPa (600 to 2500 psi) in a third embodiment.

[0075] Catalysts Employed: The slurry catalyst comprises an active catalyst in a hydrocarbon oil diluent. In one embodiment, the catalyst is a sulfided catalyst comprising at least a Group VIB metal, or at least a Group VIII metal, or at least a group IIB metal, e.g., a ferric sulfide catalyst, zinc sulfide, nickel sulfide, molybdenum sulfide, or an iron zinc sulfide catalyst. In another embodiment, the catalyst is a multi-metallic catalyst comprising at least a Group VIB metal and at least a Group VIII metal (as a promoter), wherein the metals may be in elemental form or in the form of a compound of the metal. In one example, the catalyst is a MoS₂ catalyst promoted with at least a group VIII metal compound.

[0076] In one embodiment, the catalyst is a bulk multi-metallic catalyst comprising at least one Group VIII non-noble metal and at least two Group VIB metals, and wherein the ratio of the at least two Group VIB metals to the Group VIII non-noble metal is from about 10:1 to about 1:10. In another embodiment, the catalyst is of the formula $(M^t)_a(X^u)_b(S^v)_d(C^w)_e(H^x)_f(O^y)_g(N^z)_h$, wherein M represents at least one group VIB metal, such as Mo, W, etc. or a combination thereof; and X functions as a promoter metal, representing at least one of: a non-noble Group VIII metal such as Ni, Co; a Group VIII metal such as Fe; a Group VIB metal such as Cr; a Group IVB metal such as Ti; a Group IIB metal such as Zn, and combinations thereof (X is hereinafter referred to as "Promoter Metal"). Also in the equation, t, u, v, w, x, y, z representing the total charge for each of the component (M, X, S, C, H, O and N, respectively); $ta+ub+vd+we+xf+yg+zh=0$. The subscripts ratio of b to a has a value of 0 to 5, and $(0 \leq b/a \leq 5)$. S represents sulfur with the value of the subscript d ranging from $(a + 0.5b)$ to $(5a + 2b)$. C represents carbon with subscript e having a

value of 0 to 11($a+b$). H is hydrogen with the value off ranging from 0 to 7($a+b$). O represents oxygen with the value of g ranging from 0 to 5($a + b$); and N represents nitrogen with h having a value of 0 to 0.5($a + b$). In one embodiment, subscript b has a value of 0, for a single metallic component catalyst, e.g., Mo only catalyst (no promoter).

[0077] In one embodiment, the catalyst is prepared from catalyst precursor compositions including organometallic complexes or compounds, e.g., oil soluble compounds or complexes of transition metals and organic acids. Examples of such compounds include naphthenates, pentanedionates, octoates, and acetates of Group VIB and Group VIII metals such as Mo, Co, W, etc. such as molybdenum naphthanate, vanadium naphthanate, vanadium octoate, molybdenum hexacarbonyl, and vanadium hexacarbonyl.

[0078] In one embodiment, the catalyst is a MoS_2 catalyst, promoted with at least a group VIII metal compound. In another embodiment, the catalyst is a bulk multimetallic catalyst, wherein said bulk multimetallic catalyst comprises of at least one Group VIII non-noble metal and at least two Group VIB metals and wherein the ratio of said at least two Group VIB metals to said at least one Group VIII non-noble metal is from about 10:1 to about 1:10.

[0079] In one embodiment, the catalyst feed comprises slurry catalyst having an average particle size of at least 1 micron in a hydrocarbon oil diluent. In another embodiment, the catalyst feed comprises slurry catalyst having an average particle size in the range of 1 - 20 microns. In a third embodiment, the slurry catalyst has an average particle size in the range of 2 - 10 microns. In one embodiment, the feed comprises a slurry catalyst having an average particle size ranging from colloidal (nanometer size) to about 1-2 microns. In another embodiment, the catalyst comprises catalyst molecules and/or extremely small particles that are colloidal in size (i.e., less than 100 nm, less than about 10 nm, less than about 5 nm, and less than about 1 nm). In operations, the colloidal / nanometer sized particles aggregate in a hydrocarbon diluent, forming a slurry catalyst with an average particle size in the range of 1-20 microns. In yet another embodiment, the catalyst comprises single layer MoS_2 clusters of nanometer sizes, e.g., 5-10 nm on edge.

[0080] In one embodiment, a sufficient amount of fresh catalyst and used catalyst is fed to the contacting zone(s) for each contacting zone to have a slurry (solid) catalyst concentration ranging from 2 to 30 wt. %. In a second embodiment, the (solid) catalyst concentration in the reactor ranges from 3 to 20 wt.%. In a third embodiment, from 5 to 10 wt. %.

[0081] In one embodiment, the amount of fresh catalyst feed into the contacting zone(s) range from 50 to 15000 wppm of Mo (concentration in heavy oil feed). In a second embodiment, the concentration of the fresh catalyst feed ranges from 150 to 2000 wppm Mo. In a third embodiment, from 250 to 5000 wppm Mo. In a fourth embodiment, the concentration is less than 10,000 wppm Mo. The concentration of the fresh catalyst into each contacting zone may vary depending on the contacting zone employed in the system, as catalyst may become more concentrated as volatile fractions are removed from a non-volatile resid fraction, thus requiring adjustment of the catalyst concentration.

[0082] Optional Treatment System - SDA: In one embodiment of the invention, a solvent deasphalting unit (SDA) is employed before the first contacting zone to pre-treat the heavy oil feedstock. In yet another embodiment, a solvent deasphalting unit is employed as an intermediate unit located after one of the intermediate separation zones.

[0083] SDA units are typically used in refineries to extract incremental lighter hydrocarbons from a heavy hydrocarbon stream, whereby the extracted oil is typically called deasphalted oil (DAO), while leaving a residue stream behind that is more concentrated in heavy molecules and heteroatoms, typically known as SDA Tar, SDA Bottoms, etc. The SDA can be a separate unit or a unit integrated into the upgrade system.

[0084] Various solvents may be used in the SDA, ranging from propanes to hexanes, depending on the desired level of deasphalting prior to feeding the contact zone. In one embodiment, the SDA is configured to produce a deasphalted oil (DAO) for blending with the catalyst feed or feeding directly into the contacting zones instead of, or in addition to the heavy oil feed. As such, the solvent type and operating conditions can be optimized such that a high volume and acceptable quality DAO is produced and fed to the contacting zone. In this embodiment, a suitable solvent to be used includes, but not limited to hexane or similar C_6+ solvent for a low volume SDA Tar and high volume DAO. This scheme would allow for the vast majority of the heavy oil feed to be upgraded in the subsequent contacting zone, while the very heaviest, bottom of the barrel bottoms that does not yield favorable incremental conversion economics due to the massive hydrogen addition requirement, to be used in some other manner.

[0085] In one embodiment, all of the heavy oil feed is pre-treated in the SDA and the DAO product is fed into the first contacting zone, or fed according to a split feed scheme with at least a portion going to a contacting zone other than the first in the series. In another embodiment, some of the heavy oil feed (depending on the source) is first pre-treated in the SDA and some of the feedstock is fed directly into the contacting zone(s) untreated. In yet another embodiment, the DAO is combined with the untreated heavy oil feedstock as one feed stream to the contacting zone(s). In another embodiment, the DAO and the untreated heavy oil feedstock are fed to the system as in separate feed conduits, with the DAO going to one or more of the contacting zones and the untreated heavy oil feed going to one or more of the same or different contacting zones.

[0086] In an embodiment wherein the SDA is employed as an intermediate unit, the non-volatile fraction containing the slurry catalyst and optionally minimum quantities of coke / asphaltenes, etc. from at least one of the separation zones is sent to the SDA for treatment. From the SDA unit, the DAO is sent to at least one of the contacting zones as a feed stream by itself, in combination with a heavy oil feedstock as a feed, or in combination with the bottom stream from one

of the separation zones as a feed. The DA Bottoms containing asphaltenes are sent away to recover metal in any carry-over slurry catalyst, or for applications requiring asphaltenes, e.g., blended to fuel oil, used in asphalt, or utilized in some other applications.

[0087] In one embodiment, the quality of the DAO and DA Bottoms is varied by adjusting the solvent used and the desired recovery of DAO relative to the heavy oil feed. In an optional pretreatment unit such as the SDA, the more DAO oil that is recovered, the poorer the overall quality of the DAO, and the poorer the overall quality of the DA Bottoms. With respect to the solvent selection, typically, as a lighter solvent is used for the SDA, less DAO will be produced, but the quality will be better, whereas if a heavier solvent is used, more DAO will be produced, but the quality will be lower. This is due to, among other factors, the solubility of the asphaltenes and other heavy molecules in the solvent.

[0088] Controlling Heavy Metal Deposit - Optional Water Injection: As used herein, the front-end contacting zone (or the first contacting zone) means the 1st reactor in a system with three or less contacting zones. In another embodiment of a system with more than three contacting zones, the first front-end contacting zone may include both first and second reactors. In yet another embodiment, the first contacting zone means the 1st reactor only.

[0089] As used herein, the term "water" is used to indicate either water and / or steam. In one embodiment to control heavy metal deposit, water is optionally injected into the system. In one embodiment, the injection is at a rate of about 1 to 25 wt. % (relative to the heavy oil feedstock). In one embodiment, a sufficient amount of water is injected for a water concentration in the system in the range of 2 to 15 wt. %. In a third embodiment, a sufficient amount is injected for a water concentration in the range of 4 to 10 wt. %.

[0090] The water can be added to the heavy oil feedstock before or after preheating. In one embodiment, a substantial amount of water is added to the heavy oil feedstock admixture that is to be preheated, and a substantial amount of water is added directly to the front end contacting zone(s). In another embodiment, water is added to the front-end contacting zone(s) via the heavy oil feedstock only. In yet another embodiment, at least 50% of the water is added to the heavy oil feedstock mixture to be heated, and the rest of the water is added directly to the front end contacting zone(s).

[0091] In one embodiment, the water introduced into the system at the preheating stage (prior to the preheating of the heavy oil feedstock), in an amount of about 1 to about 25 wt. % of the incoming heavy oil feedstock. In one embodiment, water is added to as part of the heavy oil feed to all of the contacting zones. In another embodiment, water is added to the heavy oil feed to the first contacting zone only. In yet another embodiment, water is added to the feed to the first two contacting zones only.

[0092] In one embodiment, water is added directly into the contacting zone at multiple points along the contacting zone, in ratio of 1 to 25 wt. % of the heavy oil feedstock. In yet another embodiment, water is added directly into the first few contacting zones in the process which are the most prone to deposits of heavy metals.

[0093] In one embodiment, some of the water is added to the process in the form of dilution steam. In one embodiment, at least 30% of the water added is in the form of steam. In the embodiments where water is added as dilution steam, the steam may be added at any point in the process. For example, it may be added to the heavy oil feedstock before or after preheating, to the catalyst / heavy oil mixture stream, and / or directly into the vapor phase of the contacting zones, or at multiple points along the first contacting zone. The dilution steam stream may comprise process steam or clean steam. The steam may be heated or superheated in a furnace prior to being fed into the upgrade process.

[0094] It is believed that the presence of the water in the process favorably alter the metallic compound sulfur molecular equilibrium, thus reducing the heavy metal deposit. In one embodiment, the addition of water is also believed to help control / maintain a desired temperature profile in the contacting zones. In yet another embodiment, it is believed that the addition of water to the front end contacting zone(s) lowers the temperature of the reactor(s). As the reactor temperature is lowered, it is believed that the rate of reaction of the most reactive vanadium species slows down, allowing vanadium deposition onto the slurry catalyst to proceed in a more controlled manner and for the catalyst to carry the vanadium deposits out of the reactor thus limiting the solid deposit in the reactor equipment.

[0095] In one embodiment, the addition of water reduces the heavy metal deposits in the reactor equipment at least 25% compared to an operation without the addition of water, for a comparable period of time in operation, e.g., for at least 2 months. In another embodiment, the addition of water reduces heavy metal deposits of at least 50% compared to an operation without the water addition. In a third embodiment, the addition of water reduces heavy metal deposits of at least 75% compared to an operation without the water addition.

[0096] Controlling Heavy Metal Deposit with Reactor Temperature: In one embodiment, instead of and / or in addition to the addition of water to the front end contacting zone(s), the temperature of the front end contacting zone(s) most prone to heavy metal deposits is lowered.

[0097] In one embodiment, the temperature of the first reactor is set to be at least 10°F. (5.56°C.) lower than the next reactor in series. In a second embodiment, the first reactor temperature is set to be at least 15°F. (8.33°C.) than the next reactor in series. In a third embodiment, the temperature is set to be at least 20°F. (11.11°C.) lower. In a fourth embodiment, the temperature is set to be at least 25°F. (13.89°C.) lower than the next reactor in series.

[0098] Controlling Heavy Metal Deposit with Recycled Catalyst Stream: In one embodiment, at least a portion of the non-volatile stream from at least one of the separation zones and / or an interstage deasphalting unit is recycled back

to the front end contacting zone(s) to control the heavy metal deposits.

[0099] According to the present invention this recycled stream ranges from 35 to 50 wt% of the total heavy oil feedstock to the process.

[0100] In one embodiment, the recycled stream comprises non-volatile materials from the last separation zone in the system, containing unconverted materials, heavier hydrocracked liquid products, slurry catalyst, small amounts of coke, asphaltenes, etc. In one embodiment, the recycled stream contains between 3 to 30 wt. % slurry catalyst. In another embodiment, the catalyst amount ranges from 5 to 20 wt. % . In yet another embodiment, the recycled stream contains 1 to 15 wt. % slurry catalyst.

[0101] In some embodiments, it is believed that with additional recycled catalyst provided by the recycled stream, more catalytic surface area (via the slurry catalyst in the recycled stream) is available to spread the heavy metal deposition, thus there is less trapping or deposition on the equipment. The additional catalyst surface areas provided by the recycled stream helps minimize overloading the catalyst surface with heavy metal deposit, leading to deposition on the process equipment (walls, internals, etc.).

[0102] Process Conditions: In one embodiment, the process condition being controlled to be more or less uniformly across the contacting zones. In another embodiment, the condition varies between the contacting zones for upgrade products with specific properties.

[0103] In one embodiment, the upgrade system is maintained under hydrocracking conditions, e.g., at a minimum temperature to effect hydrocracking of a heavy oil feedstock. In one embodiment, the system operates at a temperature ranging from 400°C (752°F) to 600°C (1112 °F), and a pressure ranging from 10 MPa (1450 psi) to 25 MPa (3625 psi). In one embodiment, the process condition being controlled to be more or less uniformly across the contacting zones. In another embodiment, the condition varies between the contacting zones for upgrade products with specific properties.

[0104] In one embodiment, the contacting zone process temperature ranges from about 400°C (752°F) to about 600°C (1112 °F), less than 500°C (932°F) in another embodiment, and greater than 425°C. (797°F) in another embodiment. In one embodiment, the system operates with a temperature difference between the inlet and outlet of a contacting zone ranging from 5 to 50°F (2.7 to 27.7°C). In a second embodiment, from 10 to 40°F (5.5 to 22.2°C).

[0105] The temperature of the separation zone is maintained within $\pm 90^\circ\text{F}$ (about $\pm 50^\circ\text{C}$) of the contacting zone temperature in one embodiment, within $\pm 70^\circ\text{F}$ (about $\pm 38.9^\circ\text{C}$) in a second embodiment, within $\pm 15^\circ\text{F}$ (about $\pm 8.3^\circ\text{C}$) in a third embodiment, and within $\pm 5^\circ\text{F}$ (about $\pm 2.8^\circ\text{C}$) in a fourth embodiment. In one embodiment, the temperature difference between the last separation zone and the immediately preceding contacting zone is within $\pm 50^\circ\text{F}$ (about $\pm 28^\circ\text{C}$).

[0106] The process pressure in the contacting zones ranges from about 10 MPa (1,450 psi) to about 25 MPa (3,625 psi) in one embodiment, about 15 MPa (2,175 psi) to about 20 MPa (2,900 psi) in a second embodiment, less than 22 MPa (3,190 psi) in a third embodiment, and more than 14 MPa (2,030 psi) in a fourth embodiment. In one embodiment, the pressure of the separation zone is maintained within ± 68.9 to ± 344.7 kPa (± 10 to ± 50 psi) of the preceding contacting zone in one embodiment, and within ± 13.7 to ± 68.9 kPa (± 2 to ± 10 psi) in a second embodiment.

[0107] In one embodiment, the upgrade system is configured for optimal operation, e.g., efficiency with much less downtime due to equipment plugging compared to the prior art with less than 689.5 kPa (100 psi) pressure drop. The optimal efficiency is obtained in one embodiment with minimal pressure drop in the system, wherein the pressure of the separation zone is maintained within ± 68.9 to ± 689.5 kPa (± 10 to ± 100 psi) of the preceding contacting zone in one embodiment, within ± 137.8 to ± 517.1 kPa (± 20 to ± 75 psi) in a second embodiment, and within ± 344.7 to ± 689.5 kPa (± 50 to ± 100 psi) in a third embodiment. As used here, the pressure drop refers to the difference between the exit pressure of the preceding contacting zone X and the entry pressure of the separation Y, with (X-Y) being less than 689.5 kPa (100 psi).

[0108] Optimal efficiency can also be obtained with minimal pressure from one contacting zone to the next contacting zone for a system operating sequentially, with the pressure drop being maintained to be 689.5 kPa (100 psi) or less in one embodiment, and 517.1 kPa (75 psi) or less in a second embodiment, and less than 344.8 kPa (75 psi) in a third embodiment. The pressure drop herein refers to the difference between the exit pressure of one contacting zone and the entry pressure of the next contacting zone.

[0109] In one embodiment, the contacting zone is in direct fluid communication to the next separation zone or contacting zone for a minimum pressure drop. As used herein, direct fluid communication means that there is free flow from the contacting zone to the next separation zone (or the next contacting zone) in series, with no flow restriction. In one embodiment, direct fluid communication is obtained with no flow restriction due to presence of valves, orifices (or a similar device), or changes in pipe diameter.

[0110] In one embodiment, the minimal pressure drop from the contacting zone to the next separation zone or contacting zone (upon entering the separating zone or the contacting zone) is due to piping components, e.g., elbows, bends, tees in the line, etc., and not due to the use of pressure reducing device such as valves, control valves, etc. to induce the pressure drop as in the prior art. In the prior art, it is taught that the separation zone functions as an interstage pressure differential separator.

[0111] In one embodiment, the minimal pressure drop is induced by friction loss, wall drag, volume increase, and changes in height as the effluent flows from the contacting zone to the next equipment in series. If valves are used in the once through system, the valves are selected / configured such that the pressure drop from one equipment, e.g., the contacting zone, to the next piece of equipment is kept to be at 100 psi or lower.

[0112] In one embodiment, the liquid hourly space velocity (LHSV) of the heavy oil feed will generally range from about 0.025 h⁻¹ to about 10 h⁻¹, about 0.5 h⁻¹ to about 7.5 h⁻¹, about 0.1 h⁻¹ to about 5 h⁻¹, about 0.75 h⁻¹ to about 1.5 h⁻¹, or about 0.2 h⁻¹ to about 10 h⁻¹. In some embodiments, LHSV is at least 0.5 h⁻¹, at least 1 h⁻¹, at least 1.5 h⁻¹, or at least 2 h⁻¹. In some embodiments, the LHSV ranges from 0.025 to 0.9 h⁻¹. In another embodiment, the LHSV ranges from 0.1 to 3 LHSV. In another embodiment, the LHSV is less than 0.5 h⁻¹.

[0113] In one embodiment wherein all of the non-volatile fractions stream from at least a separation zone is sent to the SDA unit for deasphalting, the solid deposit in the last contacting zone in the system decreases by at least 10% (in terms of deposit volume) after a similar run time compared to a prior art operation without deasphalting with the SDA unit. In a second embodiment, the solid deposit decreases by at least 20% compared to an operation without the use of the interstage SDA unit. In a third embodiment, the solid deposit decreases at least 30%.

[0114] In various embodiments, it is found that by diverting some, if not all, of the fresh catalyst to contacting zone(s) other than the first one in the system, the overall cracking efficiency of the heavy oil feedstock was not noticeably or at all impacted, as compared to the prior art feed scheme with all of the fresh catalyst going to the 1st contact zone. In one embodiment, the shift in the location of the fresh catalyst injection yields a significant boost in overall catalytic activity, with the improved quality of the non-volatile stream from the last separation zone in the system (bleed stream, "Stripper Bottoms" or STB) in terms of API, viscosity, MCR level, nickel, Hydrogen / Carbon ratio, and hot heptane asphaltenes (HHA) level. In some other embodiments, less catalyst bleeding is also observed with the overall improvement in catalytic activity.

[0115] In one embodiment, the STB product improvements include a nickel reduction of at least 10%, in a second embodiment, a nickel reduction of at least 20%. In a third embodiment, a Ni level of less than 10 ppm.

[0116] In one embodiment, the MCR reduction in the STB is at least 5%. In another embodiment, the MCR reduction is at least 10%. In a third embodiment, the MCR level is less than 13 wt. %.

[0117] In one embodiment, the STB displays an API viscosity improvement of at least 15%. In a second embodiment, an API viscosity improvement of at least 30%. In a third embodiment, an API viscosity of at least 50%, going from 2.7 to 4.5. It is observed that in some embodiments, the improvement of the API is due to overall improved catalytic activity, thus resulting in a higher H/C ratio.

[0118] In embodiments with a heavy oil split feed scheme, it is found that by diverting a portion of the heavy oil feedstock from the first contacting zone to at least one other contact zone in the series, the overall coke formation is substantially reduced as compared to the feed scheme of the prior art with all of the heavy oil feedstock going to the 1st contacting zone. Additionally, with at least a portion of the heavy oil feedstock being diverted to contacting zones other than the 1st in the system, there is some liquid dilution in these contacting zones (that would not have been present in the prior art scheme). The liquid dilution allows a more uniform catalyst concentration profile across all reactors in the system, thus protecting the last reactor against solids level excursion that could lead to operation problems.

[0119] In some embodiments with a heavy oil split feed scheme, it is also observed that the overall system efficiency improves as the conversion level in the reactors (contacting zones) increases, allowing for additional oil vaporization and corresponding decrease in liquid throughput and increase in catalyst concentration. This would essentially boost the efficiency of the system with a lower liquid throughput (or higher liquid residence time) and higher catalyst concentration. Additionally, with a secondary steady heavy oil feed rate directly into the last reactor, the last reactor is protected against upset conditions that could deprive this vessel of liquid flow. Hence, the heavy oil split feed scheme reduces or eliminates "over-conversion events" or "dry" conditions often observed in hydroprocessing reactors. In upgrade system running under "dry" conditions, insufficient liquid flow is present thus leading to solids buildup / coking, degrading flow patterns and / or hydrodynamics, degrading thermometry, loss of reaction volume, eventually compromised performance, stability and longevity of the operation.

[0120] Figures Illustrating Embodiments: Reference will be made to the figures to further illustrate embodiments of the invention.

[0121] Figure 1 is a block diagram schematically illustrating a system for upgrading heavy oil feedstock with reduced heavy metal deposits. First, a heavy oil feedstock is introduced into the first contacting zone in the system together with a slurry catalyst feed. In the figure, the slurry catalyst feed comprises a combination of fresh catalyst and recycled catalyst slurry as separate streams. Hydrogen may be introduced together with the feed in the same conduit, or optionally, as a separate feed stream. Water and / or steam may be introduced together with the feed and slurry catalyst in the same conduit or a separate feed stream. Although not shown, the mixture of water, heavy oil feed, and slurry catalyst can be preheated in a heater prior to feeding into the contacting zone. Although not shown, additional hydrocarbon oil feed, e.g., VGO, naphtha, in an amount ranging from 2 to 30 wt. % of the heavy oil feed can be optionally added as part of the feed stream to any of the contacting zones in the system.

[0122] Although not shown in the figures, the system may comprise recirculating / recycling channels and pumps for promoting the dispersion of reactants, catalyst, and heavy oil feedstock in the contacting zones, particularly with a high recirculation flow rate to the first contacting zone to induce turbulent mixing in the reactor, thus reducing heavy metal deposits. In one embodiment, a recirculating pump circulates through the loop reactor, thus maintaining a temperature difference between the reactor feed point to the exit point ranging from 0.5 to 27.8°C (1 to 50°F), and preferably between 1.1 and 13.9°C (2-25°F).

[0123] In the contacting zones under hydrocracking conditions, at least a portion of the heavy oil feedstock (higher boiling point hydrocarbons) is converted to lower boiling hydrocarbons, forming an upgraded product. The water / steam in the first contacting zone is expected to cut down on the heavy metal deposits onto the equipment. Although not illustrated, the temperature of the first contacting zone can be kept at least 2.8 - 13.9 °C (2-25 °F) lower than the temperature of the next contacting zone in series.

[0124] Upgraded material is withdrawn from the 1st contacting zone and sent to a separation zone, e.g., a hot separator, operated at a high temperature and high pressure similar to the contacting zone. The upgraded material may be alternatively introduced into one or more additional hydroprocessing reactors (not shown) for further upgrading prior to going to the hot separator. The separation zone causes or allows the separation of gas and volatile liquids from the non-volatile fractions. The gaseous and volatile liquid fractions are withdrawn from the top of the separation zone for further processing. The non-volatile (or less volatile) fraction is withdrawn from the bottom. Slurry catalyst and entrained solids, coke, hydrocarbons newly generated in the hot separator, etc., are withdrawn from the bottom of the separator and fed to the next contacting zone in the series. In one embodiment (not shown), a portion of the non-volatile stream is recycled back to one of the contacting zones preceding the separation zone, providing recycled catalyst for use in the hydroconversion reactions.

[0125] In one embodiment (as indicated by dotted lines), portions of the fresh catalyst feed and heavy oil feedstock are fed directly into contacting zones (reactors) other than the 1st contacting zone in the system. In one embodiment wherein portions of the heavy oil feedstock are fed directly into contacting zones other than the 1st contacting zone, water and / or steam is also provided to the contacting zones as a separate feed stream, or introduced together with the feed and slurry catalyst in the same conduit.

[0126] The liquid stream from the preceding separation zone is combined with optional fresh catalyst, optional additional heavy oil feed, optional hydrocarbon oil feedstock such as VGO (vacuum gas oil), and optionally recycled catalyst (not shown) as the feed stream for the next contacting zone in the series. Hydrogen may be introduced together with the feed in the same conduit, or optionally, as a separate feed stream. Upgraded materials along with slurry catalyst flow to the next separation zone in series for separation of gas and volatile liquids from the non-volatile fractions. The gaseous and volatile liquid fractions are withdrawn from the top of the separation zone, and combined with the gaseous and volatile liquid fractions from a preceding separation zone for further processing. The non-volatile (or less volatile) fraction stream is withdrawn and sent to the next contacting zone in series for the unconverted heavy oil feedstock to be upgraded.

[0127] In the last contacting zone, hydrogen is added along with the unconverted heavy oil feedstock, optional additional heavy oil feedstock, optional VGO feed, and optional fresh catalyst. Upgraded materials flow to the next separation zone along with slurry catalyst, wherein the upgraded products are removed overhead, and a portion of the non-volatile materials are recycled. In one embodiment, the recycled stream is sent to the first contacting zone, providing some of recycled catalyst for use in the hydroconversion reactions. In a second embodiment, the recycled stream is split amongst the contacting zones preceding the last contacting zone in the series.

[0128] In one embodiment, the system may optionally comprise an in-line hydrotreater (not shown) for treating the gaseous and volatile liquid fractions from the separation zones. The in-line hydrotreater in one embodiment employs conventional hydrotreating catalysts, is operated at a similarly high pressure (within 10 psig) as the rest of the upgrade system, and capable of removing sulfur, Ni, V, and other impurities from the upgraded products. In another embodiment, the in-line hydrotreater operates at a temperature within 100°F of the temperature of the contacting zones.

[0129] Figure 2 is a flow diagram of a heavy oil upgrade process with water injection. As shown, water 81 is injected into the system with the heavy oil feedstock, with the mixture being preheated in furnace before being introduced into the contacting zone. Water / steam can also be optionally injected into the system after the preheater as stream 82. In this embodiment, the fresh catalyst feed is split amongst the contacting zones. Recycle catalyst stream 17, water / heavy oil feedstock mixture, and hydrogen gas 2 are fed to the first contacting zone as feed 3.

[0130] Stream 4 comprising upgraded heavy oil feedstock exits the contacting zone R-10 flows to a separation zone 40, wherein gases (including hydrogen) and upgraded products in the form of volatile liquids are separated from the non-volatile liquid fraction 7 and removed overhead as stream 6. The non-volatile stream 7 is sent to the next contacting zone 20 in series for further upgrade. Non-volatile stream 7 contains slurry catalyst in combination with unconverted oil, and small amounts of coke and asphaltenes in some embodiments.

[0131] The upgrade process continues with the other contacting zones as shown, wherein the feed stream to contacting zone 20 comprises non-volatile fractions, hydrogen feed, optional VGO feed, and fresh catalyst feed 32. From contacting zone 20, stream 8 comprising upgraded heavy oil feedstock flows to separation zone 50, wherein upgraded products

are combined with hydrogen and removed as overhead product 9. Bottom stream 11 containing non-volatile fractions, e.g., catalyst slurry, unconverted oil containing coke and asphaltenes flow to the next contacting zone in the series 30.

[0132] In contacting zone 30, additional hydrogen containing gas 16, fresh catalyst 33, optional hydrocarbon feed such as VGO (not shown), optional untreated heavy oil feed (not shown), are added to the non-volatile stream from the preceding separation zone. From contacting zone 30, upgraded products, unconverted heavy oil, slurry catalyst, hydrogen, etc. are removed overhead as stream 12 and sent to the next separation zone 60. From the separator, overhead stream 13 containing hydrogen and upgraded products is combined with the overhead streams from preceding separation zones, and sent away for subsequent processing in another part of the system, e.g., to a high pressure separator and / or lean oil contactor and / or an in-line hydrotreater (not shown). A portion of the non-volatile stream 17 is removed as bleed-off stream 18. The rest is recycled back to at least one of the contacting zones (first contacting zone 10 as shown) as a recycled catalyst stream.

[0133] Figure 3 is a flow diagram of another embodiment of the heavy oil upgrade process, but with steam injection 91 instead of or in addition to the water injection stream 81.

[0134] Figure 4 is a flow diagram of another embodiment of the heavy oil upgrade process, with a recycled catalyst stream 19 ranging between 35 to 50 wt.% of total heavy oil feedstock to the process.

[0135] Figure 5 is a block diagram schematically illustrating another embodiment for upgrading heavy oil feedstock. First, a heavy oil feedstock is introduced into the first contacting zone in the system together with a slurry catalyst feed. Hydrogen may be introduced together with the feed in the same conduit, or optionally, as a separate feed stream. In one embodiment (not shown), optional hydrocarbon oil feedstock such as VGO (vacuum gas oil), naphtha, MCO (medium cycle oil), solvent donor, or other aromatic solvents, etc. in an amount ranging from 2 to 30 wt. % of the heavy oil feed. The additional hydrocarbon feedstock may be used to modify the concentration of metals and impurities in the system. In the contacting zones under hydrocracking conditions, at least a portion of the heavy oil feedstock (higher boiling point hydrocarbons) is converted to lower boiling hydrocarbons, forming an upgraded product.

[0136] Upgraded material is withdrawn from the 1st contacting zone and sent to a separation zone, e.g., a hot separator. The upgraded material may be alternatively introduced into one or more additional hydroprocessing reactors (not shown) for further upgrading prior to going to the hot separator. The separation zone causes or allows the separation of gas and volatile liquids from the non-volatile fractions. The gaseous and volatile liquid fractions are withdrawn from the top of the separation zone for further processing. The non-volatile (or less volatile) fraction is withdrawn from the bottom. Slurry catalyst, small amounts of heavier hydrocracked liquid products, and entrained solids, coke, hydrocarbons newly generated in the hot separator, etc., are withdrawn from the bottom of the separator and fed to the next contacting zone in the series. In one embodiment (not shown), a portion of the non-volatile stream is recycled back to the contacting zone directly preceding the separation zone, in an amount equivalent to 35 to 40 wt. % of the total heavy oil feed.

[0137] The non-volatile stream from the preceding separation zone containing unconverted feedstock is combined with additional fresh catalyst, optional additional heavy oil feed, and optionally recycled catalyst (not shown) as the feed stream for the next contacting zone in the series.

[0138] In the next contacting zone and under hydrocracking conditions, more of the heavy oil feedstock is upgraded to lower boiling hydrocarbons. Upgraded materials along with slurry catalyst flow to the next separation zone in series for separation of gas and volatile liquids from the non-volatile fractions. The non-volatile (or less volatile) stream is withdrawn from the bottom. The gaseous and volatile liquid fractions are withdrawn from the top of the separation zone (and combined with the gaseous and volatile liquid fractions from a preceding separation zone) as "upgraded" products for further processing or blending, e.g., for final blended products meeting specifications designated by refineries and / or transportation carriers.

[0139] In one embodiment (not shown), the non-volatile material containing unconverted materials is sent to the next contacting zone in series. In another embodiment as shown, the non-volatile material is recycled back to one of the contacting zones in the system, with a portion of the material being bled off for further processing, e.g., going to a solvent deasphalting unit, a catalyst deoiling unit and subsequently a metal recovery system. The recycled non-volatile material in one embodiment is an amount equivalent to 2 to 50 wt. % of the heavy oil feedstock to the system, providing recycled catalyst for use in the hydroconversion reactions.

[0140] Depending on the operating conditions, the type of catalyst fed into the contacting zone and the concentration of the slurry catalyst, in one embodiment, the outlet stream from the contacting zones comprises a ratio of 20:80 to 60:40 of upgraded products to unconverted heavy oil feed. In one embodiment, the amount of upgraded products out of the first contacting zone is in the range of 30-35% to unconverted heavy oil product of 65-70%.

[0141] Although not shown in the figures, the system may optionally comprise recirculating / recycling channels and pumps for promoting the dispersion of reactants, catalyst, and heavy oil feedstock in the contacting zones. In one embodiment, a recirculating pump circulates through the loop reactor a volumetric recirculation ratio of 5:1 to 15:1 (recirculated amount to heavy oil feed ratio), thus maintaining a temperature difference between the reactor feed point to the exit point ranging from 5.5 to 27.7 °C (10 to 55°F), and preferably between 11.1 to 22.2 (20 to 40°F).

[0142] In one embodiment, the system may optionally comprise an in-line hydrotreater (not shown) for treating the

gaseous and volatile liquid fractions from the separation zones. The in-line hydrotreater in one embodiment employs conventional hydrotreating catalysts, is operated at a similarly high pressure (within 68.9 kPa (10 psig) in one embodiment, and 344.8 kPa (50psig) in a second embodiment) as the rest of the upgrade system, and capable of removing sulfur, Ni, V, and other impurities from the upgraded products.

[0143] Figure 6 is a block diagram schematically illustrating another embodiment of an upgrade system, wherein a solvent deasphalting unit is employed for pre-treating some, if not all of the heavy oil feed to the system. The de-asphaltened oil (DAO) can be fed directly to the contacting zone(s) or combined with a heavy oil feed stream as a feedstock. In some embodiment, other hydrocarbon materials, e.g., VGO, can also be combined with the heavy oil feed and / or the DAO as the feedstock for some of the contacting zone(s). All of the fresh catalyst can be fed directly to the 1st contacting zone in the system, or diverted to other contacting zone(s) in the series.

[0144] Figure 7 is a flow diagram of a heavy oil upgrade process with a fresh catalyst split feed scheme, wherein some of the fresh catalyst feed is diverted from the first reactor to other reactors in the process. As shown, the fresh catalyst feed is split amongst the various contacting zones as feed streams 31, 32, and 33. Fresh catalyst feed 31 is combined with the recycle catalyst stream 17 and fed to the first contacting zone as slurry catalyst feed 3. Hydrogen gas 2 and heavy oil feedstock 1 are combined with slurry catalyst 3 as feed into the first contacting zone 10. In this embodiment, heavy oil feedstock is preheated in furnace 80 before being introduced into the contacting zone as heated oil feed 4.

[0145] Stream 5 comprising upgraded heavy oil feedstock exits the contacting zone 10 and flows to a separation zone 40, wherein gases (including hydrogen) and volatile upgraded products are separated from the non-volatile fractions 7 and removed overhead as stream 6. The non-volatile fractions stream 7 is sent to the next contacting zone 20 in series for further upgrade. Stream 7 contains slurry catalyst in combination with unconverted oil, and small amounts of coke and asphaltenes in some embodiments.

[0146] The upgrade process continues with the other contacting zones as shown, wherein stream 7 is combined with hydrogen feed 15 and fresh catalyst 32 as feed stream into contacting zone 20. Although not shown, the streams can also be fed to the contacting zone in separate conduits. Stream 8 comprising upgraded heavy oil feedstock flows to separation zone 50, wherein upgrade products are combined with hydrogen and removed as overhead product 9. Bottom stream 11 containing catalyst slurry, unconverted oil (plus small amounts of coke and asphaltenes in some embodiments) is combined with a fresh catalyst stream 33 and a fresh supply of hydrogen 16 as feed stream to the next contacting zone 30. Stream 12 exits the contacting zone and flows to separation zone 60, wherein upgraded products and hydrogen are removed overhead as stream 13. Some of the bottom stream 17 from the separation zone, which contains catalyst slurry, unconverted oil plus small amounts of coke and asphaltenes in some embodiments, is recycled back to the 1st contacting zone 10 as recycled stream 19. The rest of the bottom stream 17 is removed as bleed-off stream 18 and sent to other processes in the system for catalyst de-oiling, metal recovery, etc.,. Although not shown, vapor stream 14 containing the upgraded products and hydrogen in one embodiment is subsequently processed in another part of the system, e.g., in a high pressure separator and / or lean oil contactor.

[0147] Figure 8 illustrates another embodiment of the invention, wherein reactors having internal separators are employed, thus separate hot separators / flash drums are not necessary for phase separation. In this upgrade system, a reactor differential pressure control system (not shown) is employed, regulating the product stream out of the top of each reactor-separator. External pumps (not shown) may be employed to aid in the dispersion of the slurry catalyst in the system and help control the temperature in the system.

[0148] In the embodiment of Figure 8 as shown, all of fresh catalyst is diverted to the 2nd and 3rd contacting zones in the system. Recycled catalyst stream 19 provides slurry catalyst feed to the first contacting zone, and optionally, to other contacting zone(s) in the system. Also as shown, additional hydrocarbon oil feed, e.g., VGO, naphtha, in an amount ranging from 2 to 30 wt. % of the heavy oil feed can be optionally added as part of the feed stream to any of the contacting zones in the system.

[0149] Figure 9 illustrates an embodiment of the invention wherein all of the fresh catalyst feed 99 is fed directly to the last contacting zone in the upgrade system, with other contacting zone(s) in the system simply getting a portion of the recycled catalyst stream 19.

[0150] Figure 10 illustrates is an embodiment of a heavy oil split feed scheme. As shown, some of heavy oil feed is diverted from the 1st reactor and fed directly to the 2nd contacting zone in the system as heavy oil feed stream 42. Also as shown, recycled catalyst is optionally sent to the 2nd contacting zone in the system along with portions of the fresh catalyst as stream 32.

[0151] The following examples are given as non-limitative illustration of aspects of the present invention.

[0152] Comparative Example 1: Heavy oil upgrade experiments were carried out in a pilot system having three gas-liquid slurry phase reactors connected in series with three hot separators, each being connected in series with the reactors. The upgrade system was run continuously for about 50 days.

[0153] A fresh slurry catalyst used was prepared according to the teaching of US Patent No. 2006/0058174, i.e., a Mo compound was first mixed with aqueous ammonia forming an aqueous Mo compound mixture, sulfided with hydrogen compound, promoted with a Ni compound, then transformed in a hydrocarbon oil (other than heavy oil feedstock) at a

temperature of at least 176.7°C (350°F) and a pressure of at least 1379 kPa (200 psig), forming an active slurry catalyst to send to the first reactor.

[0154] The hydroprocessing conditions were as follows: a reactor temperature (in three reactors) of about 825°F (440.5°C); a total pressure in the range of 16547.4 kPa to 17.926.4 Kpa (2400 to 2600 psig); a fresh Mo/ fresh heavy oil feed ratio (wt. %) 0.20 - 0.40; fresh Mo catalyst /total Mo catalyst ratio 0.125 - 0.250; total feed LHSV about 0.070 to 0.15; and H₂ gas rate 7500 to 20000 SCF/ bbl (1335.7 to 3561.9 1./1.).

[0155] Effluent taken from each reactor was sent to the separator (connected in series), and separated into a hot vapor stream and a non-volatile stream. Vapor streams were removed from the top of the high pressure separators and collected for further analysis ("HPO" or high-pressure overhead streams). The non-volatile stream containing slurry catalyst and unconverted heavy oil feedstock was removed from the separator and sent to the next reactor in series.

[0156] A portion of the non-volatile stream from the last separator in an amount of 30 wt. % of heavy oil feedstock was recycled (STB), and the rest was removed as a bleed stream (in an amount of about 15 wt. % of the heavy oil feedstock). The STB stream contains about 10 to 15 wt. % slurry catalyst.

[0157] The feed blend to the system was high metals heavy crude with the properties specified in Table 1.

Table 1

	VR feed
API gravity at 60/60	-
Specific gravity	1.0760
Sulfur (wt %)	5.27015
Nitrogen (ppm)	7750
Nickel (ppm)	135.25
Vanadium (ppm)	682.15
Carbon (wt %)	83.69
Hydrogen (wt %)	9.12
H/C Ratio	0.109

[0158] After 50 days of operation, the operation was shut down. The reactor, distributor and internal thermowell were visually inspected. All three pieces show significant built-up of deposit, with approximately 28.5% of the volume of the front-end (1st) reactor being lost due to deposits of heavy metals. Analysis of the used slurry catalyst in the bleed stream over the 50 day period showed an increasing deficit in vanadium, suggesting that the deposit build up inside the front end reactor was not only happening but actually worsening over the course of the run. The performance of the process also suffered, due to the loss in the reaction volume.

[0159] Example 2: Example 1 was repeated, except that the temperature of the 1st reactor was decreased 11.1 °C (20°F) from about 440.5 to 429.4°C (825°F to 805°F), the recycled catalyst rate was increased from 30 wt. % (in Example 1) to about 40 wt. % of the heavy oil feed rate, and water was added to the front end reactor at a rate equivalent to 5 wt. % of the heavy oil feed rate. The system ran for 54 days before shutdown.

[0160] Water injection was carried out by adding water to the fresh catalyst, then the water catalyst mixture was added to an autoclave along with the heavy oil feed and hydrogen, with the mixture being pre-heated to a temperature of about 350°F (176.7°C).

[0161] Analysis of the used slurry catalyst in the bleed stream over the 54 day period showed a fairly close agreement between the amount of vanadium expected to exit the process and the amount of vanadium in the catalyst in the bleed stream, suggesting that vanadium trapping has significantly reduced, thus heavy metal deposit in the equipment.

[0162] The analytical results were further confirmed by visual inspections of the reactor internals, distributor, and internal thermowell. The equipment was significantly cleaner in Example 2, with only 6.6% of the front end reactor volume being lost due to heavy metal deposits.

[0163] Comparative Example 3: Heavy oil upgrade experiments were carried out in a pilot system having three gas-liquid slurry phase reactors connected in series with two hot separators. The hot separators are connected in series with the 1st and 3rd reactors respectively, with no hot separator following the 2nd reactor. The gas-liquid slurry phase reactors were continuously stirred reactors. The upgrade system was run continuously for about 70 days.

[0164] A fresh slurry catalyst used was prepared according to the teaching of US Patent No. 2006/0058174, i.e., a Mo compound was first mixed with aqueous ammonia forming an aqueous Mo compound mixture, sulfided with hydrogen/sulfur compound, promoted with a Ni compound, then transformed in a hydrocarbon oil (other than heavy oil feedstock) at a temperature of at least 176.6°C(350°F) and a pressure of at least 1379 kPa (200 psig), forming an active slurry catalyst.

[0165] In Comparative Example 3, all of the fresh catalyst slurry was sent to the first reactor in the system, for a

concentration of fresh slurry catalyst in heavy oil ranging from 2,000 to 5,000 ppm, expressed as weight of metal (molybdenum) to weight of heavy oil feed. The hydroprocessing conditions were as follows: a reactor temperature of 435 - 440.5 °C (815 - 825°F); a total pressure in the range of 16547.4 to 17926.4 kPa (2400 to 2600 psig); a fresh Mo/ fresh heavy oil feed ratio (wt. %) 0.20 - 0.40; fresh Mo catalyst /total Mo catalyst ratio 0.1; total feed LHSV 0.10 to 0.15; and H₂ gas rate 1780.9 to 2671.4 l./l. (10000 to 15000 SCF/bbl).

[0166] Effluent taken from the 1st and 3rd reactors was introduced into the hot separators connected in series with the reactors, and separated into a hot vapor stream and a non-volatile stream. Vapor streams were removed from the top of the high pressure separators and collected for further analysis ("HPO" or high-pressure overhead streams). The non-volatile stream containing slurry catalyst and unconverted heavy oil feedstock was removed from the bottom of the 1st separator and sent to the 2nd reactor in series. Effluent from the 2nd reactor was sent directly to the 3rd reactor as feedstock.

[0167] A portion of the non-volatile stream from the last separator in an amount of 5 - 15 wt. % of heavy oil feedstock was removed as the bleed-off stream, for an overall conversion rate of 98 to 98.5% of heavy oil feed to distillate products. The rest of the non-volatile stream, the "Stripper Bottoms product" or STB, containing the bulk of the catalyst (in an amount of 80 to 95% of total slurry catalyst entering the system) was recycled back to the first reactor for maintaining the flow of catalyst through the upgrade system. The STB stream contains about 7 to 20 wt% slurry catalyst. The STB was also analyzed to evaluate the overall performance of the system.

[0168] The feed blend to the system was a heavy oil feed with the properties specified in Table 2.

Table 2

		VR Properties
	API gravity at 60/60	4.6
	Specific gravity	1.04
	Sulfur (wt %)	1.48
	Nitrogen (ppm)	11069
	Nickel (ppm)	118.8
	Vanadium (ppm)	108.7
	Carbon (wt %)	83.57
	Hydrogen (wt %)	10.04
	MCR (wt %)	20.7
	Viscosity @ 100°C. (cSt)	20796
	Pentane Asphaltenes (wt %)	13.9
	Fraction Boiling above 1000°F (537.8°C) in wt% 100%	

[0169] Example 4: After 70 days with all of the fresh catalyst to the 1st reactor (in Comparative Example 3), the location of fresh catalyst supply was shifted from the 1st to the 3rd reactor, with the first two reactors relying entirely on recycled catalyst feed stream for 28 days. All other process conditions remained the same. HPO and STB products were collected, analyzed, and compared with the results of Comparative Example 3. There was no significant change in HPO product quality. With respect to the STB product, the results are as follows:

Table 3

STB Product properties	Comparative Example 3	Example 4
Wt %VR (BP 537.8°C=1000°F)	15.9	15.3
Wt % HVGO (BP 426.7°C=800°F)	49.8	48.6
Wt % VGO (BP 343.3°C=650°F)	79.8	80.0
API	2.7	4.5
Sulfur (wt. %)	0.12	0.16
Nitrogen (ppm)	12711	12335
MCR (wt. %)	14.7	12.4
Hydrogen / Carbon ratio	0.098	0.102
Ni (ppm)	10.8	7.9
Hot heptane asphaltenes, ppm	174255	119713

(continued)

STB Product properties	Comparative Example 3	Example 4
Viscosity @70°C, cSt	68.4	47.3

[0170] The results show that diverting the fresh catalyst to the last contacting zone in the system did not trigger changes in product nitrogen levels. However, there was a change in the sulfur level, which could be due to the unusually low sulfur level in the heavy oil feed to the system and the high sulfur level in the VGO oil used in the slurry catalyst feed. It is therefore possible that injecting the fresh catalyst into the last reactor penalized the product sulfur by providing less time for the VGO oil carrier (in the slurry catalyst) to react, resulting into a higher product sulfur level. It is further noted that by diverting the fresh catalyst to the last reactor yielded a STB product with improved properties, including API, viscosity, MCR, HHA, nickel, and H/C ratio. The improvement in STB product API did not correlate with an improvement in the distillation of the STB product. In other words, the STB product API did not improve due to additional cracking in a lighter product distillation, but due to improved catalytic activity, resulting into a higher H/C ratio.

[0171] With respect to the system operation in the 28-day run, there was no evidence of pressure-drop buildup or plugging around the front end reactors to suggest any coking or solid build-ups. There was no measurable negative impact on the overall conversion rate. The results suggest the used catalyst has retained sufficient hydrogenation activity to starve off coking, even in the presence of fresh / untreated heavy oil feedstock, indicating that a fresh catalyst split scheme still suppresses coking adequately.

[0172] Example 5: Comparative Example 3 is repeated except that 20% of the heavy oil feedstock is diverted from the 1st reactor to the 3rd reactor while other process conditions remain the same.

[0173] In comparing process stability, reactor performance, and reactor conditions between the examples, it is believed that in Comparative Example 3, the 3rd reactor has a lower liquid throughput (with no heavy oil feed) and higher catalyst concentration which are directionally beneficial for conversion purposes. However, these conditions also tend to make the last reactor more susceptible to operation upsets leading to insufficient liquid flow-through, and consequentially, more solids build-up, degrading thermometry and shortening of the process run-time.

[0174] In Example 5 with a portion of the heavy oil feedstock being fed directly to the last reactor, it is anticipated that the preceding reactors (1st and 2nd) with a decrease in liquid throughput (as a portion of the heavy oil feedstock is diverted) and a corresponding increase in catalyst concentration will operate more efficiently and with a higher conversion rate. Additionally, with more liquid dilution in the 3rd reactor, there is a more uniform catalyst concentration profile across all three reactors.

[0175] It is further anticipated that as the last reactor in the series gets a portion of the heavy oil feed, dry conditions associated with insufficient liquid flow is obviated. As the last reactor is protected from over-conversion events or dry conditions, there is less solid build-up or coke deposition. It is also expected that the last reactor is less susceptible to operation upsets, e.g., wide swings in temperature, pressure, flows, etc.

[0176] For the purpose of this specification and appended claims, unless otherwise indicated, all numbers expressing quantities, percentages or proportions, and other numerical values used in the specification and claims, are to be understood as being modified in all instances by the term "about." Accordingly, unless indicated to the contrary, the numerical parameters set forth in the following specification and attached claims are approximations that may vary depending upon the desired properties sought to be obtained and / or the precision of an instrument for measuring the value, thus including the standard deviation of error for the device or method being employed to determine the value. The use of the term "or" in the claims is used to mean "and/or" unless explicitly indicated to refer to alternatives only or the alternative are mutually exclusive, although the disclosure supports a definition that refers to only alternatives and "and/or." The use of the word "a" or "an" when used in conjunction with the term "comprising" in the claims and/or the specification may mean "one," but it is also consistent with the meaning of "one or more," "at least one," and "one or more than one." Furthermore, all ranges disclosed herein are inclusive of the endpoints and are independently combinable. In general, unless otherwise indicated, singular elements may be in the plural and vice versa with no loss of generality. As used herein, the term "include" and its grammatical variants are intended to be non-limiting, such that recitation of items in a list is not to the exclusion of other like items that can be substituted or added to the listed items.

[0177] It is contemplated that any aspect of the invention discussed in the context of one embodiment of the invention may be implemented or applied with respect to any other embodiment of the invention. Likewise, any composition of the invention may be the result or may be used in any method or process of the invention. This written description uses examples to disclose the invention, including the best mode, and also to enable any person skilled in the art to make and use the invention. The patentable scope is defined by the claims, and may include other examples that occur to those skilled in the art. Such other examples are intended to be within the scope of the claims if they have structural elements that do not differ from the literal language of the claims, or if they include equivalent structural elements with insubstantial differences from the literal languages of the claims. All citations referred herein are expressly incorporated

herein by reference.

Claims

1. A process for hydroprocessing a heavy oil feedstock, the process employing a plurality of contacting zones and separation zones, wherein the process comprises a plurality of slurry reactors for contacting zones, the process comprising:

providing a heavy oil feedstock, a hydrogen containing gas, and a slurry catalyst;
combining the heavy oil feedstock, the hydrogen containing gas, and the slurry catalyst in a first contacting zone under hydrocracking conditions to convert at least a portion of the heavy oil feedstock to lower boiling hydrocarbons, forming upgraded products;

sending a mixture comprising the upgraded products, the slurry catalyst, the hydrogen containing gas, and unconverted heavy oil feedstock to a first separation zone, whereby the upgraded products are removed with the hydrogen containing gas from the first separation zone as a first overhead stream, and the slurry catalyst, heavier hydrocracked liquid products and unconverted heavy oil feedstock are removed from the first separation zone as a first non-volatile stream;

sending the first non-volatile stream to a contacting zone other than the first contacting zone, which is maintained under hydrocracking conditions with additional hydrogen containing gas feed to convert at least a portion of the heavy oil feedstock to lower boiling hydrocarbons, forming additional upgraded products;

sending a mixture comprising the additional upgraded products, slurry catalyst, hydrogen containing gas, and unconverted heavy oil feedstock to a separation zone other than the first separation zone, whereby the upgraded products are removed with hydrogen containing gas as an overhead stream, and the slurry catalyst and the unconverted heavy oil feedstock are removed as a second non-volatile stream; and

wherein the slurry catalyst to the first contacting zone comprises at least a portion of the non-volatile stream from one of the separation zones as a recycled catalyst stream, and wherein the recycled catalyst stream is between 35 to 50 wt. % of the heavy oil feedstock;

wherein the first contacting zone is operated at a temperature of at least 10°F (5.56°C) lower than the next contacting zone.

Patentansprüche

1. Verfahren zum Hydroprocessing eines Schweröl-Einsatzstoffs, bei dem mehrere Kontaktzonen und Trennzonen eingesetzt werden, wobei das Verfahren mehrere Slurry-Reaktoren für Kontaktzonen umfasst, wobei das Verfahren Folgendes umfasst:

Bereitstellen eines Schweröl-Einsatzstoffs, eines wasserstoffhaltigen Gases und eines Slurry-Katalysators;
Vereinigen des Schweröl-Einsatzstoffs, des wasserstoffhaltigen Gases und des Slurry-Katalysators in einer ersten Kontaktzone unter Hydrocracking-Bedingungen zur Umwandlung mindestens eines Teils des Schweröl-Einsatzstoffs in niedriger siedende Kohlenwasserstoffe unter Bildung von veredelten Produkten;

Schicken eines Gemischs, das die veredelten Produkte, den Slurry-Katalysator, das wasserstoffhaltige Gas und nicht umgesetzten Schweröl-Einsatzstoff umfasst, zu einer ersten Trennzone, wodurch die veredelten Produkte mit dem wasserstoffhaltigen Gas aus der ersten Trennzone als erster Kopfstrom ausgetragen werden und der Slurry-Katalysator, schwerere hydrogecrackte flüssige Produkte und nicht umgesetzter Schweröl-Einsatzstoff aus der ersten Trennzone als ein erster nichtflüchtiger Strom ausgetragen werden;

Schicken des ersten nichtflüchtigen Stroms zu einer von der ersten Kontaktzone verschiedenen Kontaktzone, die unter zusätzlicher Zufuhr von wasserstoffhaltigem Gas unter Hydrocracking-Bedingungen gehalten wird, zur Umwandlung mindestens eines Teils des Schweröl-Einsatzstoffs in niedriger siedende Kohlenwasserstoffe unter Bildung von zusätzlichen veredelten Produkten;

Schicken eines Gemischs, das die zusätzlichen veredelten Produkte, Slurry-Katalysator, wasserstoffhaltiges Gas und nicht umgesetzten Schweröl-Einsatzstoff umfasst, zu einer von der ersten Trennzone verschiedenen zweiten Trennzone, wodurch die veredelten Produkte mit wasserstoffhaltigem Gas als Kopfstrom ausgetragen werden und der Slurry-Katalysator und der nicht umgesetzte Schweröl-Einsatzstoff als ein zweiter nichtflüchtiger Strom ausgetragen werden; und

wobei der Slurry-Katalysator für die erste Kontaktzone mindestens einen Teil des nichtflüchtigen Stroms aus einer der Trennzonen als rezyklierten Katalysatorstrom umfasst und wobei der rezyklierte Katalysatorstrom

zwischen 35 bis 50 Gew.-% des Schweröl-Einsatzstoffs ausmacht;
wobei die erste Kontaktzone bei einer Temperatur betrieben wird, die mindestens 10 °F (5,56 °C) niedriger ist
als die der nächsten Kontaktzone.

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Revendications

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1. Procédé d'hydrotraitement d'une charge d'huile lourde, le procédé utilisant une pluralité de zones de contact et de zones de séparation, dans lequel le procédé comprend une pluralité de réacteurs à boues liquides pour les zones de contact, le procédé comprenant :

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la fourniture d'une charge d'huile lourde, d'un gaz contenant de l'hydrogène et d'un catalyseur en suspension ;
la combinaison de la charge d'huile lourde, du gaz contenant de l'hydrogène et du catalyseur en suspension
dans une première zone de contact dans des conditions d'hydrocraquage pour convertir au moins une partie
de la charge d'huile lourde en hydrocarbures à point d'ébullition plus bas, formant ainsi des produits valorisés ;
l'envoi d'un mélange comprenant les produits valorisés, le catalyseur en suspension, le gaz contenant de
l'hydrogène et la charge d'huile lourde non convertie vers une première zone de séparation, moyennant quoi
les produits valorisés sont éliminés avec le gaz contenant de l'hydrogène de la première zone de séparation
sous la forme d'un premier courant de produit de tête, et le catalyseur en suspension, les produits liquides
hydro-craqués plus lourds et la charge d'huile lourde non convertie sont éliminés de la première zone de
séparation sous la forme d'un premier courant non volatil ;

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l'envoi du premier courant non volatil vers une zone de contact autre que la première zone de contact, qui est
maintenue dans des conditions d'hydrocraquage avec une alimentation supplémentaire en gaz contenant de
l'hydrogène pour convertir au moins une partie de la charge d'huile lourde en hydrocarbures à point d'ébullition
plus bas, formant ainsi des produits valorisés supplémentaires ;

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l'envoi d'un mélange comprenant les produits valorisés supplémentaires, le catalyseur en suspension, le gaz
contenant de l'hydrogène et la charge d'huile lourde non convertie vers une zone de séparation autre que la
première zone de séparation, moyennant quoi les produits valorisés sont éliminés avec le gaz contenant de
l'hydrogène sous la forme d'un courant de produit de tête, et le catalyseur en suspension et la charge d'huile
lourde non convertie sont éliminés sous la forme d'un second courant non volatil ; et

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dans lequel le catalyseur en suspension vers la première zone de contact comprend au moins une partie du
courant non volatil provenant de l'une des zones de séparation sous la forme d'un courant de catalyseur recyclé,
et dans lequel le courant de catalyseur recyclé représente entre 35 et 50 % en poids de la charge d'huile lourde ;
dans lequel la première zone de contact est utilisée à une température inférieure d'au moins 10 °F (5,56 °C) à
celle de la zone de contact suivante.

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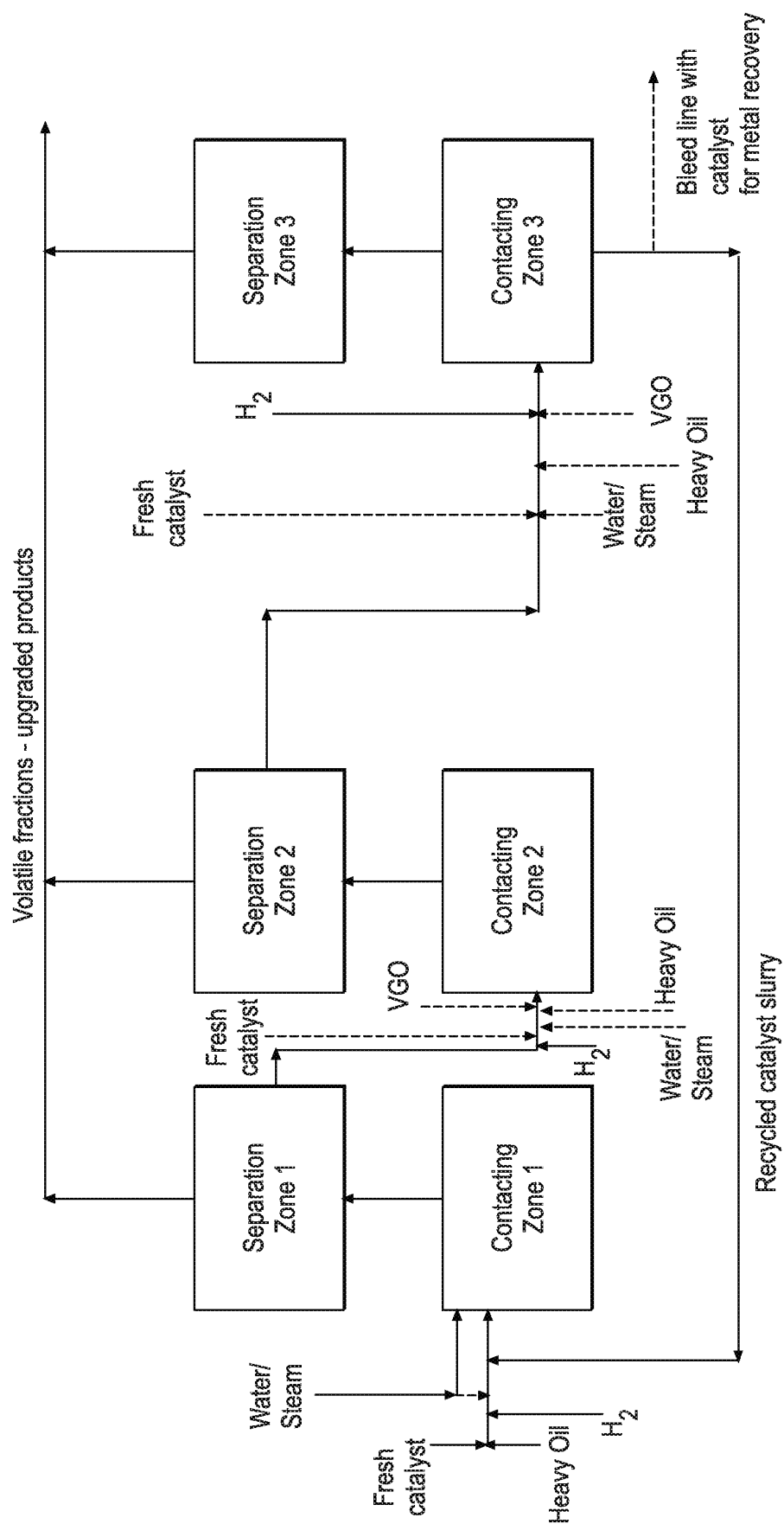


FIG. 1

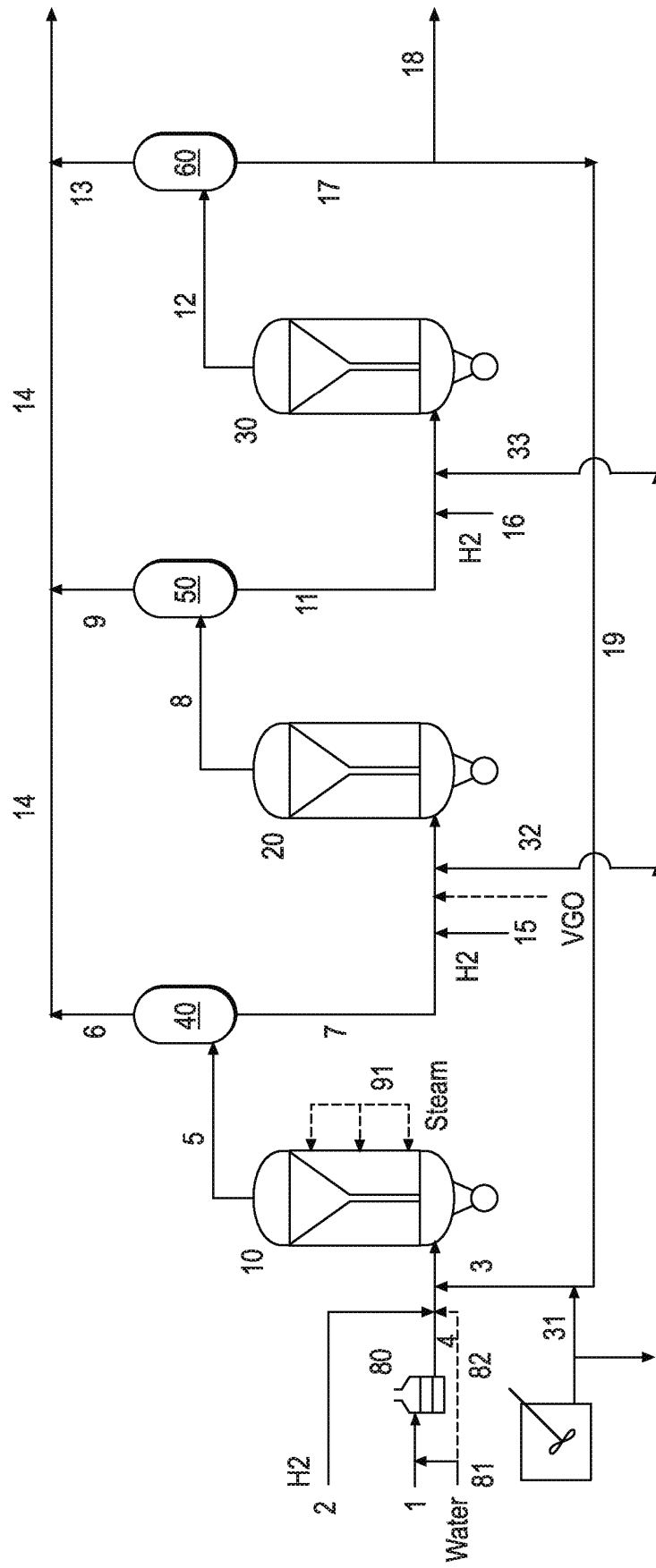


FIG. 2

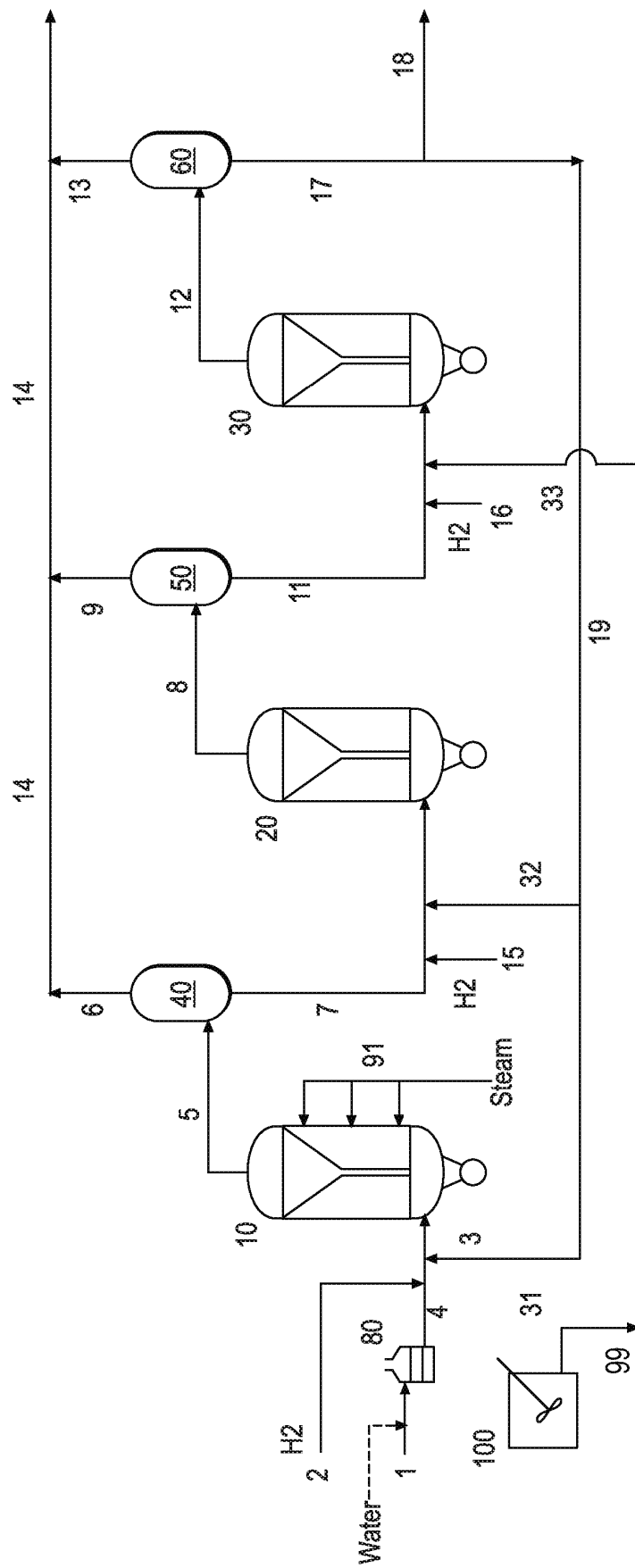


FIG. 3

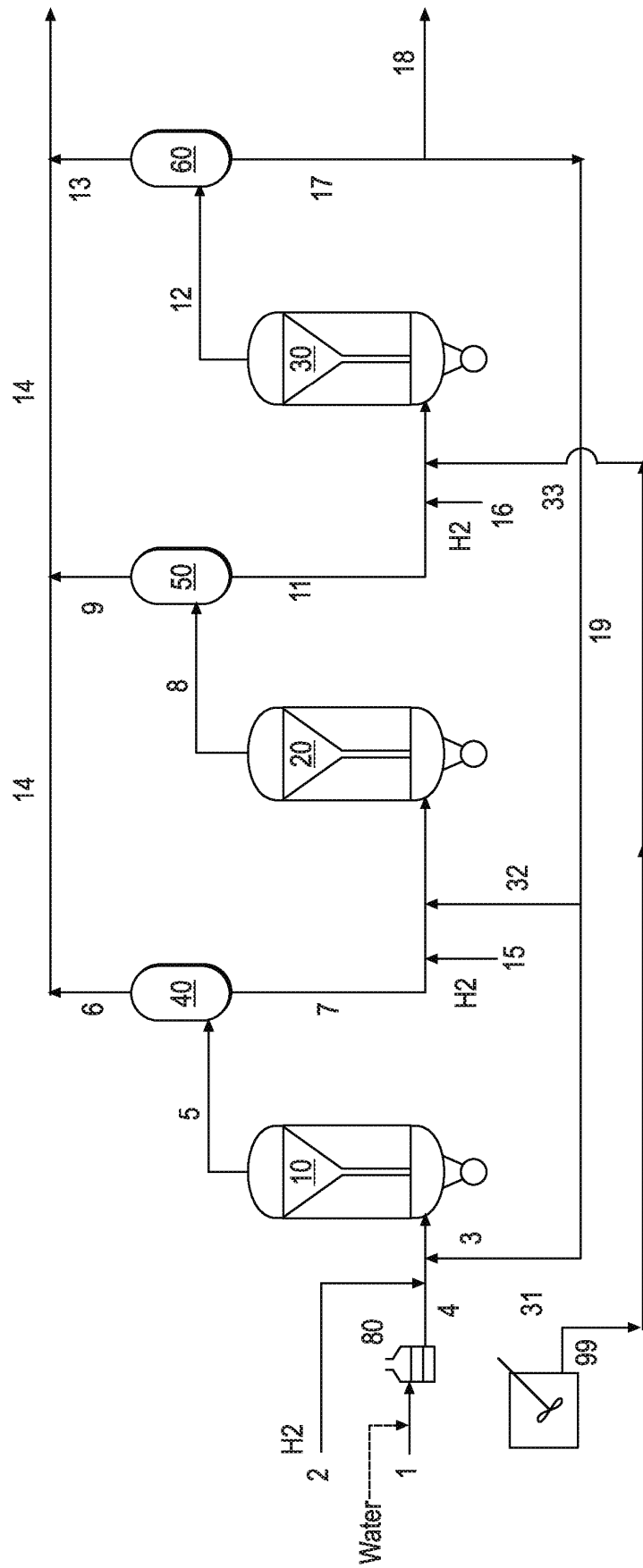


FIG. 4

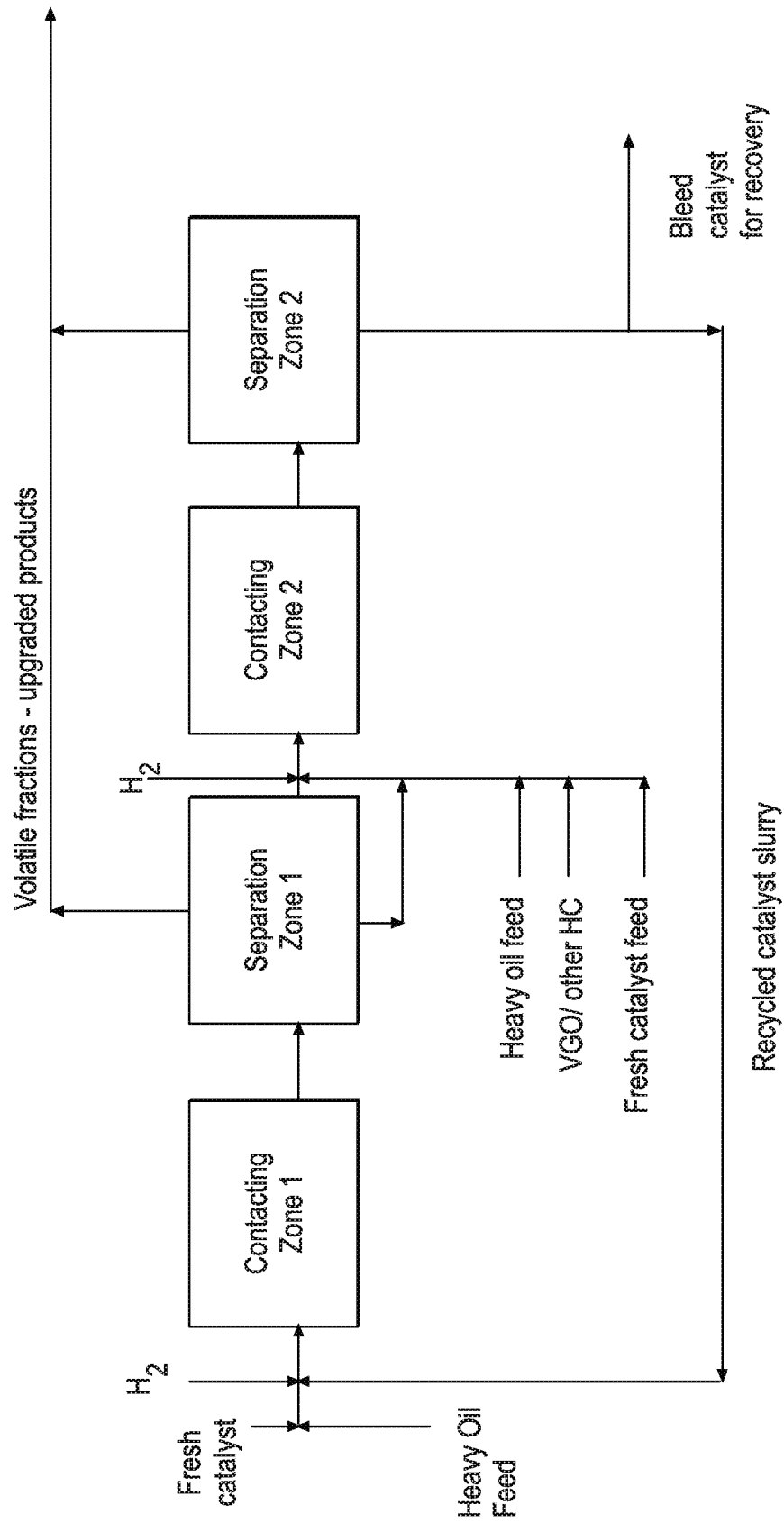
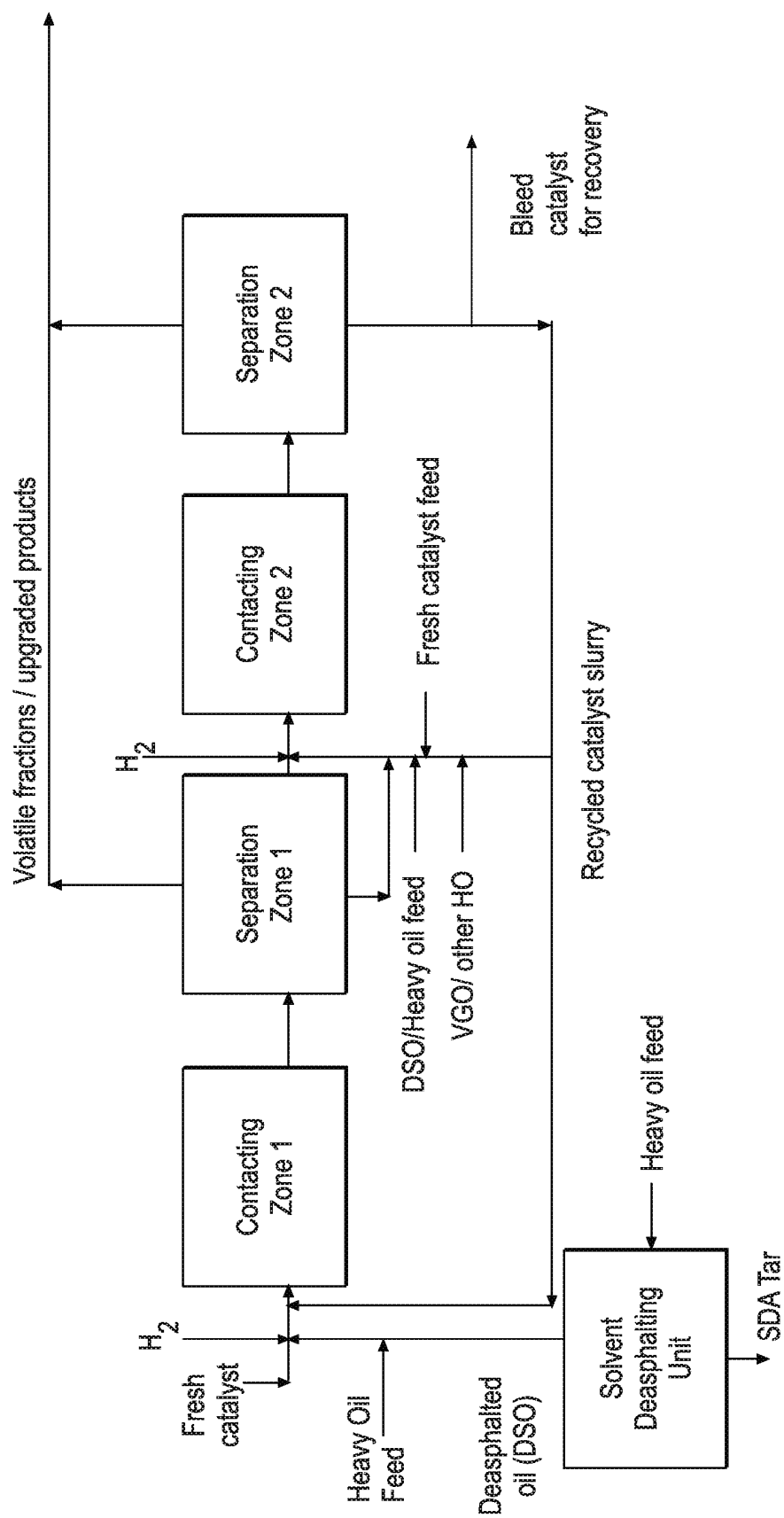


FIG. 5



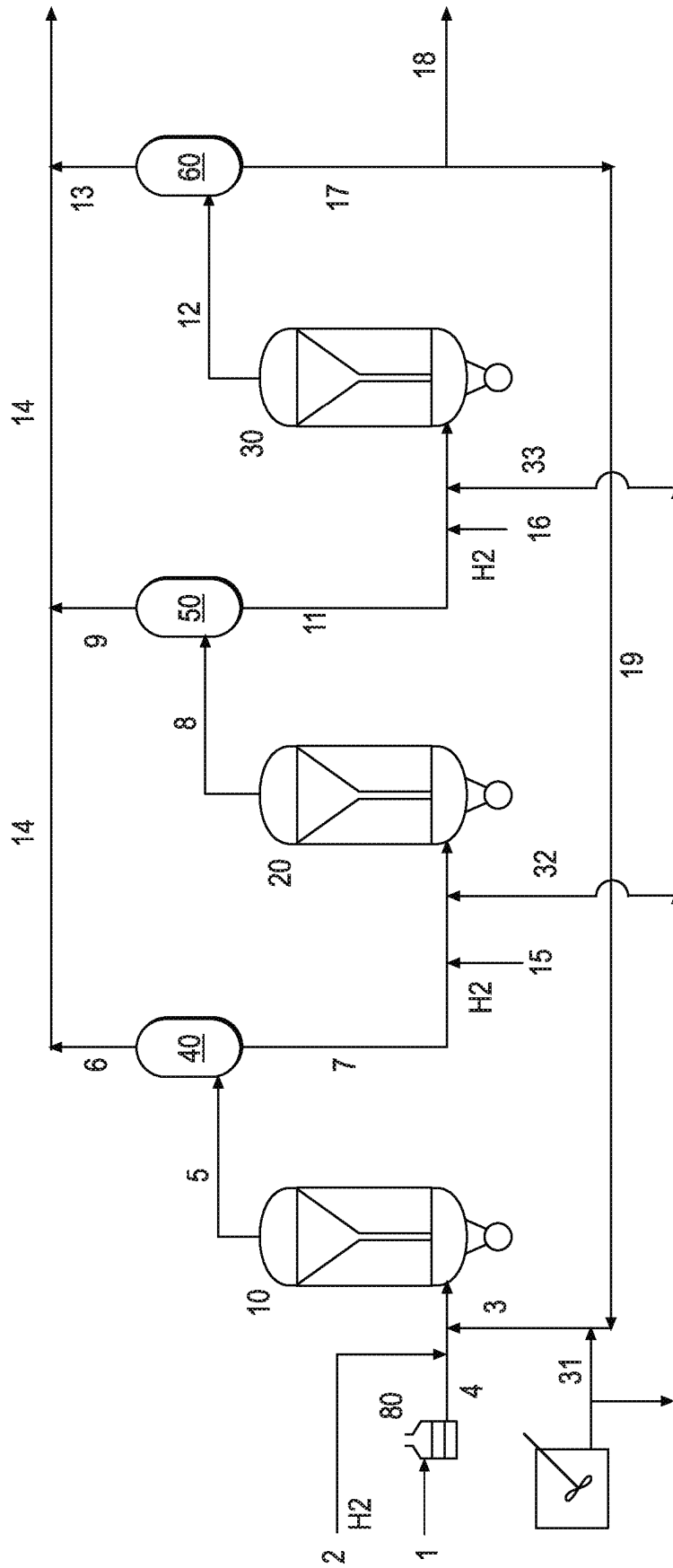


FIG. 7

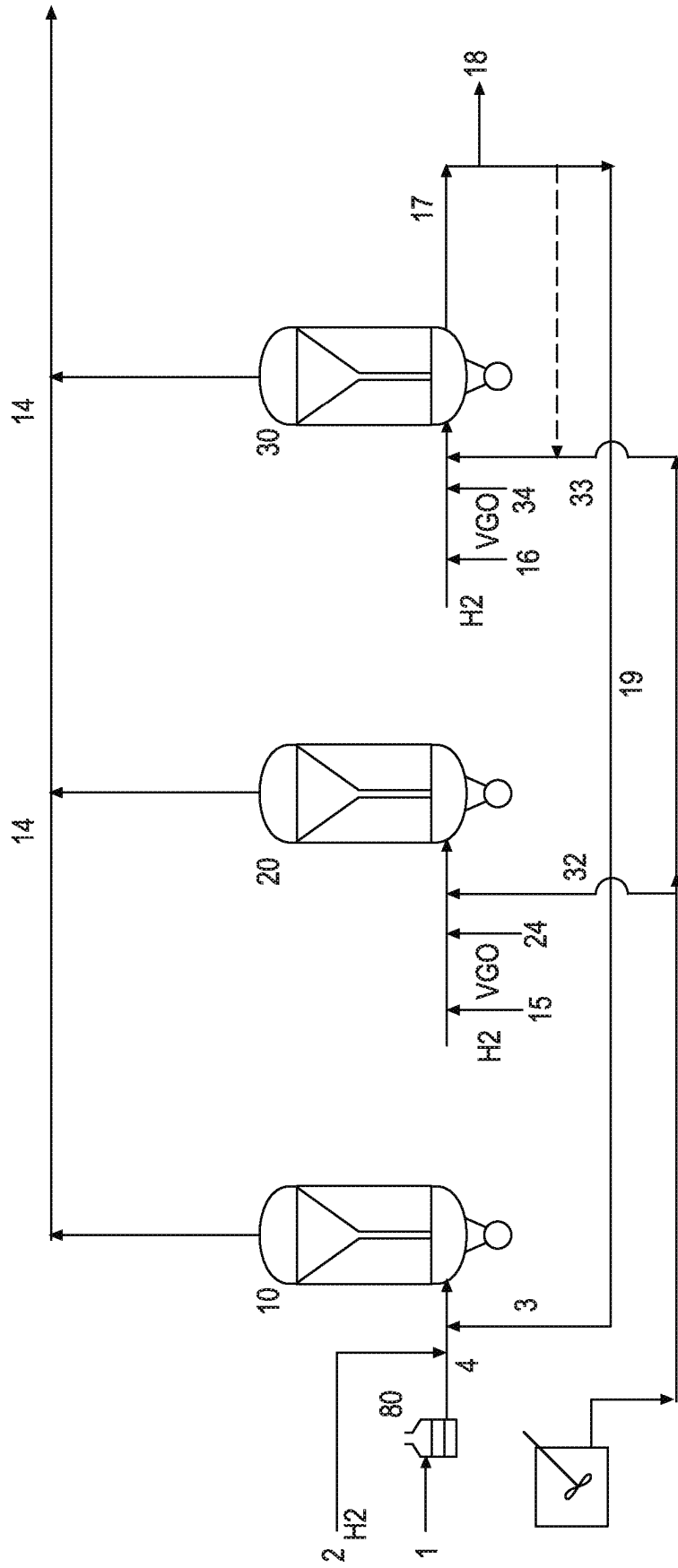
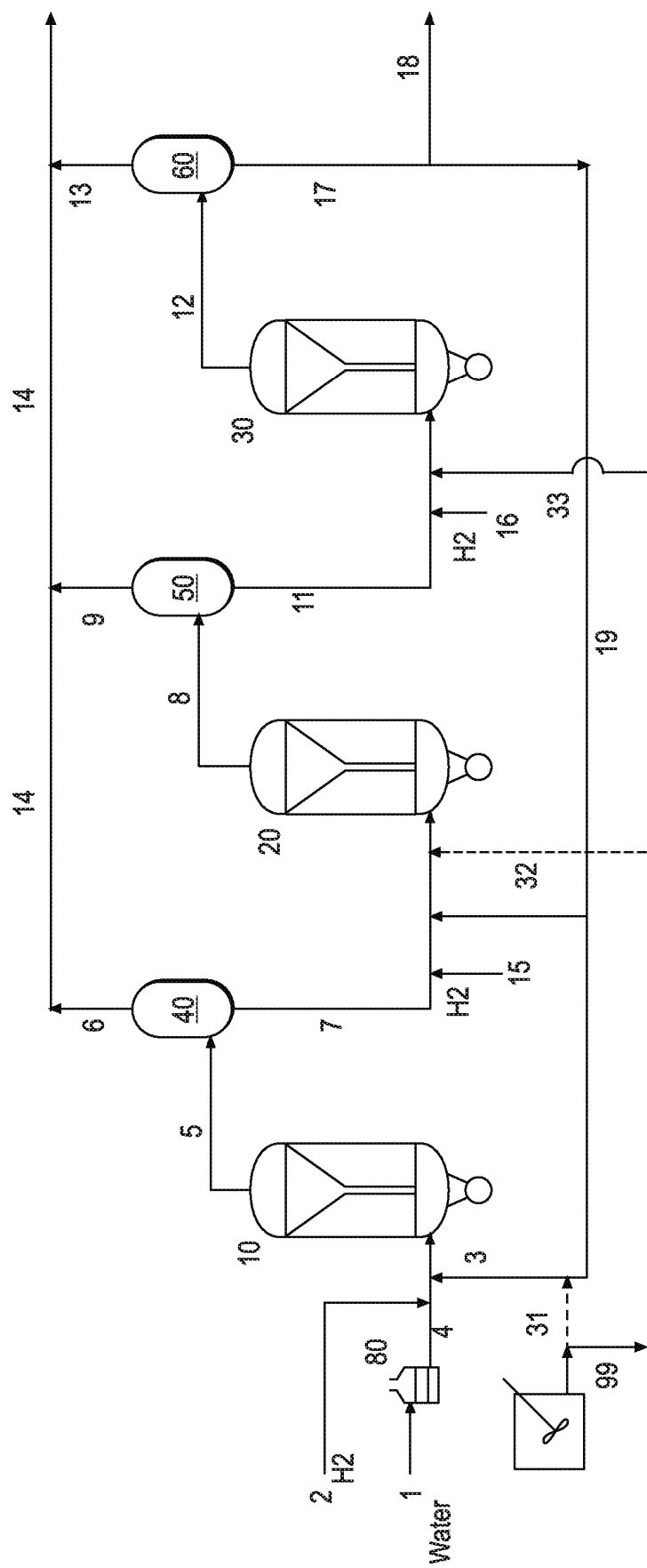


FIG. 8



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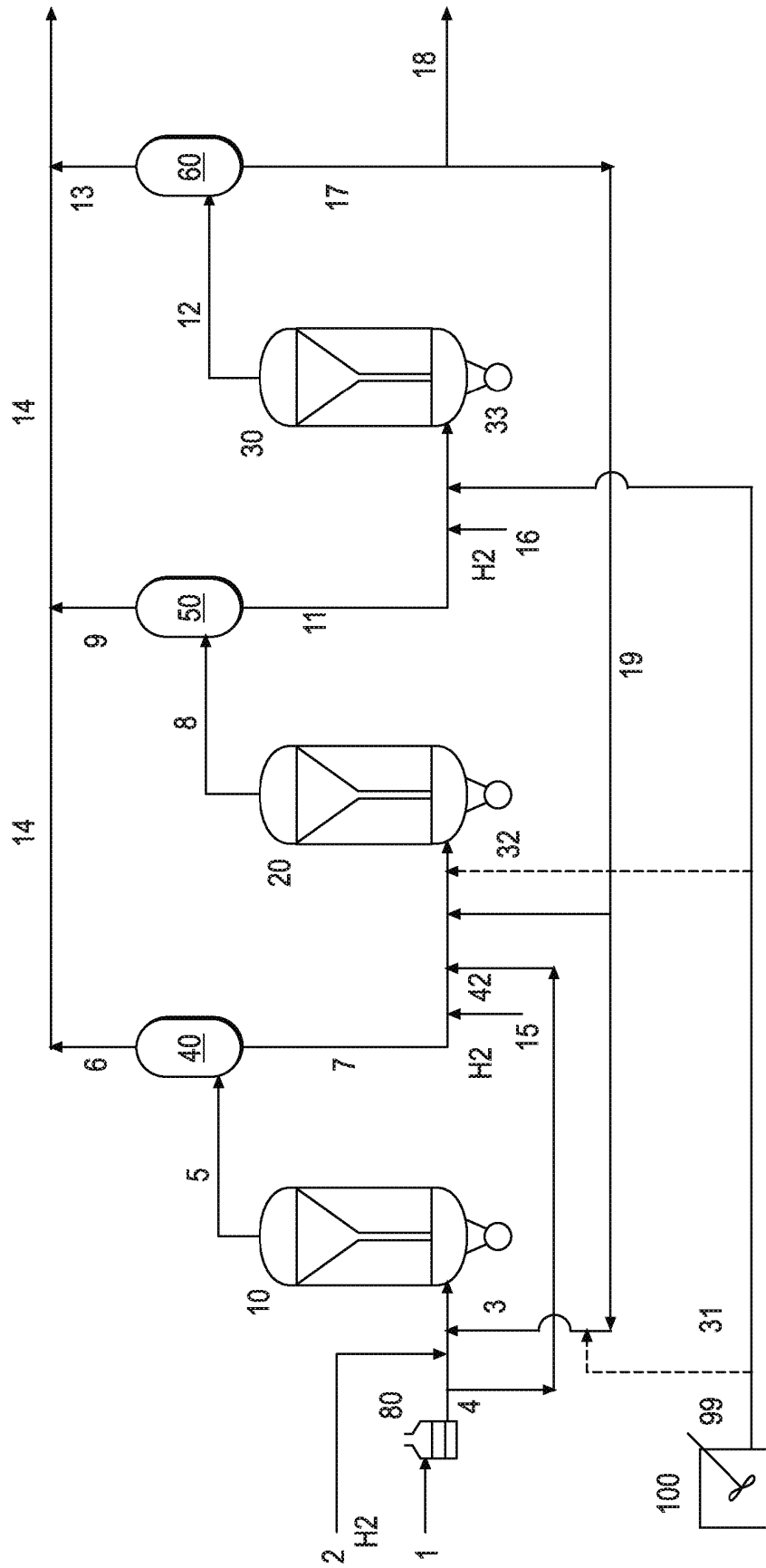


FIG. 10

REFERENCES CITED IN THE DESCRIPTION

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