The present invention pertains to a method for hydroprocessing a heavy hydrocarbon oil, comprising bringing a heavy hydrocarbon oil in a first stage into contact with hydroprocessing catalyst I in the presence of hydrogen, after which the effluent of the first stage is contacted in whole or in part with hydroprocessing catalyst II in the presence of hydrogen. The method of the invention combines efficient contaminant removal with high residue conversion and low sediment formation. The invention also comprises a combination of catalysts for use in the above method.
TWO-STAGE HPC PROCESS
CROSS REFERENCE TO RELATED APPLICATIONS

Priority of this application is based on Japanese Patent Application No. 2001-174072, filed Jun. 8, 2001, and PCT Application Serial No. PCT/EP02/06664, the contents of which are incorporated by reference herein.

BACKGROUND OF THE INVENTION

1. Field of the Invention

2. Prior Art

Hydrocarbon oils containing 50 wt. % or more of components with a boiling point of 538°C or higher are called heavy hydrocarbon oils. These include atmospheric residue (AR) and vacuum residue (VR), which are produced in petroleum refining. It is desired to remove impurities such as sulfur from these heavy hydrocarbon oils by hydroprocessing, and to convert them into lighter oils, which have a higher economic value.

The hydroprocessing of heavy hydrocarbon oils is done in ebullating bed operation or in fixed bed operation.

For ebullating bed operation, various catalysts have been proposed. Generally, these catalysts are capable of efficiently removing sulfur, Conradson carbon residue (CCR), various metals, nitrogen and/or asphaltene. However, it was found that the decomposition of asphaltene, an aggregate of condensed aromatic compounds which is in good balance with the rest of the feedstock, is generally accompanied by the formation of sediment and sludge. Sediment can be determined by the Shell hot filtration solid test (SHFST). (see Van Kerkenoot et al., J. Inst. Pet., 37, pp. 596-604 (1951)). Its ordinary content is said to be about 0.19 to 1 wt. % in product with a boiling point of 340°C or higher collected from the bottom of a flash drum.

Sediment formed during hydroprocessing may settle and deposit in such apparatuses as heat exchangers and reactors, and because it threatens to close off the passage, it can seriously hamper the operation of these apparatuses. Especially in the hydroprocessing of heavy hydrocarbon feeds containing large amounts of vacuum residue, sediment formation is an important factor, and there is therefore need for a process for effecting efficient contaminant removal in combination with low sediment formation and high conversion.

Japanese patent No. 765055 discloses a process for hydroprocessing a heavy hydrocarbon oil containing sulfur impurities and metallic impurities at least in two stages. In the first stage, a catalyst containing 0.1 to 5 wt. % of metal oxide was used to effect hydrometallization and hydrocracking. In the second stage, a catalyst containing 7 to 30 wt. % of a metal oxide was used to effect hydrodesulfurization. However, due to, among other features, the high metals content of the second catalyst, asphaltene conversion in this process is not adequate.

Japanese patent No. 8325580 A discloses a catalytic hydroconversion method for a heavy feedstock. In the first stage a catalyst with a carrier material selected from alumina, silica, and a combination thereof which comprises 2 to 25 wt. % of oxides of active metals selected from cadmium, chromium, cobalt, iron, molybdenum, nickel, tin, tungsten or a combination thereof, is applied at a reaction temperature of 438 to 468°C, a hydrogen partial pressure of 105 to 245 kg/cm², and a volume hourly space velocity of 0.3 to 1.0 h⁻¹. In the second stage a similar catalyst is used at a reaction temperature of 371 to 427°C, a hydrogen partial pressure of 105 to 245 kg/cm², and a space velocity of 0.1 to 0.8 h⁻¹. The first stage is operated at a higher temperature and a lower catalytic activity than the second stage to improve hydroconversion. However, it appears that the high temperature applied in the first stage leads to thermal condensation of asphaltene on the one hand and, on the other, the petroleum resin fractions produced with the thermal cracking of oil are unstable. This can lead to undesired cohesion and precipitation of asphaltene and coke formation in the second stage.

Japanese patent publication 653875 describes a multistage catalytic method for the high conversion of a heavy hydrocarbon liquid feedstock. In the first stage, a fixed bed or ebullated bed reactor is operated at a reaction temperature of 415 to 455°C, a hydrogen partial pressure of 70 to 211 kg/cm², and a space velocity of 0.2 to 2.0 h⁻¹. In the second stage, an ebullated bed reactor is operated under similar conditions. The catalyst used in the above method comprises a catalyst material selected from alumina, silica, and a combination thereof, and the oxide of an active metal selected from cadmium, chromium, cobalt, iron, molybdenum, nickel, tin, tungsten, and mixtures thereof. In this reference, vacuum bottoms are recirculated to achieve a high cracking rate, but the problem of asphaltene cohesion accompanying the high cracking rate operation is not solved.

U.S. Pat. No. 5,322,829 describes a two-stage ebulliating bed operation wherein the first stage catalyst may comprise 0.1-5 wt. % of nickel, calculated as oxide, and 1-15 wt. % of molybdenum, calculated as oxide, the catalyst having a surface area of at least 150 m²/g and a total pore volume of 0.8-1.4 ml/g. The second stage catalyst is not described in detail. It appears that this process is insufficient in the removal of metals and asphaltene, and shows high sediment formation.

U.S. Pat. No. 4,657,664 describes a multi-stage ebulliating bed process for the conversion of heavy hydrocarbon feeds, wherein the first stage catalyst contains 8-8 wt. % of Group VI metal oxide. The second catalyst has a high surface area the main part of which is present in pores with a low diameter. It appears that this catalyst system does not give sufficient activity and selectivity in the hydroprocessing of heavy hydrocarbon feeds.

In summary, the processes of the prior art are inadequate in inhibiting sediment formation in the hydroprocessing of heavy oil to effect hydrodesulfurization and conversion.

The object of this invention is to provide an effective process for the hydroprocessing of a heavy hydrocarbon oil containing a large amount of impurities such as sulfur, Conradson carbon residue, metals, nitrogen, and asphaltene, for adequately removing the impurities. In addition to efficient contaminant removal, the process should show low sediment formation, high asphaltene removal, and high conversion.
SUMMARY OF THE INVENTION

[0016] In one embodiment, the present invention comprises a method for hydropyroprocessing a heavy hydrocarbon oil, comprising bringing a heavy hydrocarbon oil in a first stage into contact with hydropyroprocessing catalyst I in the presence of hydrogen, after which the effluent of the first stage is contacted in whole or in part with hydropyroprocessing catalyst II in the presence of hydrogen, wherein

[0017] catalyst I comprises 7 to 20 wt. % of a Group VII B metal component, calculated as trioxide on the weight of the catalyst, and 0.5 to 6 wt. % of a Group VIII metal component, calculated as oxide on the weight of the catalyst, on a porous inorganic carrier, said catalyst having a specific surface area of at least 100 m²/g, a total pore volume of at least 0.55 ml/g, at least 50% of the total pore volume in pores with a diameter of at least 20 nm (200 Å) and at least 65% of the total pore volume in pores with a diameter of 10-120 nm (100-1200 Å), and

[0018] catalyst II comprises 7 to 20 wt. % of a Group VII B metal component, calculated as trioxide on the weight of the catalyst, and 0.5 to 6 wt. % of a Group VIII metal component, calculated as oxide on the weight of the catalyst, on a porous inorganic carrier, said catalyst having a specific surface area of at least 100 m²/g, a total pore volume of at least 0.55 ml/g, 30-80% of the pore volume in pores with a diameter of 10-20 nm (100-200 Å), and at least 5% of the pore volume in pores with a diameter of at least 100 nm (1000 Å). In Catalyst I has a larger percentage of its pore volume in pores with a diameter of at least 20 nm (200 Å) than catalyst II.

[0019] In a second embodiment, the present invention comprises a catalyst combination suitable for use in the above process, wherein the catalyst combination comprises catalysts I and II, as defined above.

[0020] Other embodiments of the invention comprise details concerning catalyst compositions and method steps, all of which are hereinafter discussed.

DETAILED DESCRIPTION OF THE INVENTION

[0021] The present invention comprises a method in which a combination of two catalysts is used to obtain advantageous effects in the hydropyroprocessing of heavy hydrocarbon oils. The present invention also relates to a combination of catalysts suitable for use in such a method.

[0022] More particularly, the present invention relates to a process suitable for the hydropyroprocessing of heavy hydrocarbon oils containing a large amount of impurities such as sulfur, metals, and asphaltene to effect hydrosulfurization (HDS), hydrodenitrogenation (HDN), asphaltene reduction (HDAI) and/or conversion into lighter products, while limiting the amount of sediment produced. The feed may also contain other contaminants such as Conradson carbon residue (CCR) and nitrogen, and carbon residue reduction (HDCCR) and hydrodenitrogenation (HDN) may also be desired processes.

[0023] The inventors have found a catalytic hydropyroprocessing method for a heavy oil based on two-stage catalytic cracking, in which both catalysts meet specific requirements as to metals content and pore size distribution. The catalyst used in the first stage is specifically designed to decrease the impurities in the heavy hydrocarbon oil. In particular, it achieves efficient asphaltene removal in combination with hydrometallization.

[0024] The second stage catalyst is tailored to effect advanced desulfurization and hydrogenation reactions while inhibiting sediment formation due to asphaltene precipitation, to allow stable operation.

[0025] The combination of the two different catalysts leads to a synergistic effect resulting in a process showing stable operation, high hydrosulfurization and conversion activity, and low sediment formation.

[0026] The catalysts used in the process according to the invention comprise catalytic materials on a porous carrier. The catalytic materials present on the catalyst according to the invention comprise a Group VII B metal and a Group VIII metal of the Periodic Table of Elements applied by Chemical Abstract Services (CAS system). The Group VIII metal used in the invention is at least one selected from nickel, cobalt, and iron. In view of performance and economy, cobalt and nickel are preferred. Nickel is especially preferred. As the Group VII B metals which can be used, molybdenum, tungsten, and chromium may be mentioned, but in view of performance and economy, molybdenum is preferred. The combination of molybdenum and nickel is particularly preferred for the catalytic materials of the catalyst according to the invention.

[0027] Based on the weight (100 wt. %) of the final catalyst, the amounts of the respective catalytic materials in the catalysts used in the process according to the invention are as follows.

[0028] The catalysts comprise 7-20 wt. %, preferably 8-16 wt. %, of Group VII B metal, calculated as trioxide. If less than 7 wt. % is used, the activity of the catalyst is insufficient. On the other hand, if more than 16 wt. % is used, the catalytic performance is not improved further.

[0029] The catalysts comprise 0.5-6 wt. %, preferably 1-5 wt. %, of Group VIII metal, calculated as oxide. If the amount is less than 0.5 wt. %, the activity of the catalysts will be too low. If more than 6 wt. % is present, the catalytic performance will not be improved further.

[0030] The total pore volume of Catalyst I and Catalyst II is at least 0.55 ml/g, preferably at least 0.6 ml/g. It is preferred if it is at most 1.0 ml/g, more preferably at most 0.9 ml/g. The determination of the total pore volume and the pore size distribution is effected via mercury penetration at a contact angle of 140° with a surface tension of 480 dynes/cm, using, for example, a mercury porosimeter Autopore II (trade name) produced by Micrometrics.

[0031] Catalyst I has a specific surface area of at least 100 m²/g. For the catalyst to meet the required pore size distribution ranges it is preferred for it to have a surface area of 100-180 m²/g, preferably 130-170 m²/g. If the surface area is less than 100 m²/g, the catalytic activity will be too low. In the present specification the surface area is determined in accordance with the BET method based on N₂ adsorption.
[0032] Catalyst I has at least 50% of the total pore volume in pores with a diameter of at least 20 nm (200 Å), preferably at least 60%. The percentage of pore volume in this range is preferably at most 80%. If the percentage of pore volume in this range is below 50%, the catalytic performance, especially the asphalene cracking activity, decreases. As a result thereof, sediment formation will increase.

[0033] Catalyst I has at least 65% of the total pore volume in pores with a diameter of 10-120 nm (100-1200 Å), preferably at least 75%, more preferably at least 85%. If the percentage of pore volume in this range is insufficient, sediment formation will increase and the residual cracking rate will decrease.

[0034] Additionally, it is preferred for Catalyst I to have less than 25% of its pore volume in pores with a diameter of 10 nm (100 Å) or less. If the percentage of pore volume present in this range is above this value, sediment formation may increase due to increased hydrogenation of the non-asphalentic feed constituents.

[0035] Catalyst I is based on a porous inorganic oxide carrier which generally comprises the conventional oxides, e.g., alumina, silica, silica-alumina, alumina with silica-alumina dispersed therein, silica-coated alumina, magnesia, zirconia, boria, and titania, as well as mixtures of these oxides. It is preferred for the carrier to consist for at least 80% of alumina, more preferably at least 90%, still more preferably at least 95%. A carrier consisting essentially of alumina is most preferred, the wording “consisting essentially of” being intended to mean that minor amounts of other components may be present, as long as they do not affect the catalytic activity of the catalyst.

[0036] Catalyst II has a surface area of at least 100 m²/g, preferably at least 150 m²/g, more preferably at least 185 m²/g. The surface area is preferably at most 250 m²/g. If the surface area is outside the claimed range, the catalytic performance is insufficient.

[0037] Catalyst II has 30-80% of the total pore volume in pores with a diameter of 10-20 nm (100-200 Å), preferably 35-75%, more preferably 40-65%. If less than 30% of the pore volume is present in this range, the catalytic performance is insufficient. If more than 80% of the total pore volume is present in this range, the sediment formation increases.

[0038] At least 5% of the total pore volume of Catalyst II is present in pores with a diameter of at least 100 nm (1000 Å), preferably between 8 and 30%, more preferably between 8 and 25%. If less than 5% of the pore volume is present in this range, the asphalene cracking activity decreases, which leads to increased sediment formation. If the percentage of pore volume present in pores with a diameter of at least 100 nm (1000 Å) is above 25%, particularly above 30%, the sediment formation may increase.

[0039] Additionally, it is preferred for Catalyst II to have less than 25% of its pore volume in pores with a diameter of 10 nm (100 Å) or less. If the percentage of pore volume present in this range is above this value, sediment formation may increase due to increased hydrogenation of the non-asphalentic feed constituents.

[0040] Catalyst II preferably has less than 50% of its pore volume present in pores with a diameter of at least 20 nm (200 Å), more preferably less than 40%. If this requirement is met, catalyst II will show particularly good hydrodesulfurization activity and good hydrogenation activity.

[0041] Catalyst II generally has a substantial portion of its surface area present in relatively large pores, that is, pores with a diameter of 15-120 nm (150-1200 Å). Preferably, the catalyst has at least 25 m²/g of surface area in pores within this range (as determined via mercury intrusion), more preferably at least 35 m²/g. On the other hand, catalyst II generally has only a limited amount of its surface area in relatively small pores, that is, pores with a diameter of 5-7 nm (50-70 Å). Preferably, the catalyst has less than 40 m²/g of surface area in pores within this range, more preferably less than 20 m²/g.

[0042] Catalyst II is also based on a porous inorganic oxide carrier which generally comprises the conventional oxides, e.g., alumina, silica, silica-alumina, alumina with silica-alumina dispersed therein, silica-coated alumina, magnesia, zirconia, boria, and titania, as well as mixtures of these oxides. It is preferred for the carrier of catalyst II to comprise at least 3.5 wt. % of silica, calculated on the weight of the final catalyst, preferably 3.5-30 wt. %, more preferably 4.12 wt. %, even more preferably 4.5-10 wt. %. If less than 3.5 wt. % of silica is present, the performance of the catalyst is less pronounced. The balance of the carrier of the Catalyst II is generally made up of alumina, optionally containing other refractory oxides, such as titania, zirconia, etc. It is preferred that the balance of the carrier of catalyst II is made up of at least 90% of alumina, more preferably at least 95%. It is preferred for the carrier of the catalyst of the invention to consist essentially of silica and alumina, the wording “consists essentially of” being intended to mean that minor amounts of other components may be present, as long as they do not affect the catalytic activity of the catalyst.

[0043] In a further preferred embodiment, Catalyst II comprises a Group IA metal component. Sodium and potassium may be mentioned as suitable materials. Sodium is preferred for reasons of performance and economy. The amount of Group IA metal is 0.1-2 wt. %, preferably 0.2-1 wt. %, more preferably 0.1-0.5 wt. %, calculated as oxide on the catalyst. If less than 0.1 wt. % is present, the desired effect will not be obtained. If more than 2 wt. % is present, or sometimes more than 1 wt. %, the activity of the catalyst will be adversely affected.

[0044] Catalyst I has a higher percentage of its pore volume present in pores with a diameter of at least 20 nm (200 Å) than catalyst II. If this requirement is not met, the advantageous effects of the present invention will not be obtained. Preferably, the % PV≥200 Å for catalyst I is at least 5% higher than the % PV≥200 Å for catalyst II, more preferably at least 10%, still more preferably at least 20%.

[0045] The catalyst particles may have the shapes and dimensions common to the art. Thus, the particles may be spherical, cylindrical, or polylobal and their diameter may range from 0.5 to 10 mm. Particles with a diameter of 0.5-3 mm, preferably 0.7-1.2 mm, for example 0.9-1 mm, and a length of 2-10 mm, for example 2.5-4.5 mm, are preferred. For use in fixed bed operation polylobal particles are preferred, because they lead to a reduced pressure drop in hydrodemonetization operations. Cylindrical particles are preferred for use in ebullating bed operations.

[0046] The carrier to be used in the catalysts to be used in the process according to the invention can be prepared by
processes known in the art. A typical production method for a carrier comprising alumina is coprecipitation of sodium aluminate and aluminum sulfate. The resulting gel is dried, extruded, and calcined, to obtain an alumina-containing carrier. Optionally, other components such as silica may be added before, during, or after precipitation.

By way of example, a process for preparing an alumina gel is described below. At first, a tank containing tap water or warm water is charged with an alkali solution of sodium aluminate, aluminum hydroxide or sodium hydroxide, etc., and an acidic aluminum solution of aluminum sulfate or aluminum nitrate, etc. is added for mixing. The hydroxide ion concentration (pH) of the mixed solution changes with the progress of the reaction. It is preferable that when the addition of the acidic aluminum solution is completed, the pH is 7 to 9, and that during mixing, the temperature is kept at a temperature for, in general, 0.5-1.5 hours, preferably for 40-80 minutes.

By way of a further example, a process for preparing a silica-containing alumina gel is described below. First, an alkali solution such as sodium aluminate, ammonium hydroxide or sodium hydroxide is fed into a tank containing tap water or hot water, an acidic solution of an aluminum source, e.g., aluminum sulfate or aluminum nitrate, is added, and the resulting mixture is mixed. The pH of the mixture changes as the reaction progresses. Preferable, after all the acid aluminum compound solution has been added, the pH is 7 to 9. After completion of the mixing an alumina hydrogel can be obtained. Then, an alkali metal silicate such as a water glass or an organic silica solution is added as silica source. To mix the silica source, it can be fed into the tank together with the acid aluminum compound solution or after the alumina hydrogel has been produced. The silica-containing alumina carrier can, for another example, be produced by combining a silica source such as sodium silicate with an alumina source such as sodium aluminate or aluminum sulfate, or by mixing an alumina gel with a silica gel, followed by moulding, drying, and calcining. The carrier can also be produced by causing alumina to precipitate in the presence of silica in order to form an aggregate mixture of silica and alumina. Examples of such methods are adding a sodium aluminate solution to a silica hydrogel and increasing the pH by the addition of, e.g., sodium hydroxide to precipitate alumina, and coprecipitating sodium silicate with aluminum sulfate. A further possibility is to immerse the alumina carrier, before or after calcination, into an impregnation solution comprising a silicon source dissolved therein.

In a following stage, the gel separated from the solution, and any commercially used washing treatment, for example a washing treatment using tap water or hot water, is carried out to remove impurities, mainly salts, from the gel. Then, the gel is shaped into particles in a manner known in the art, e.g., by way of extrusion, beadling or pelletizing.

Finally, the shaped particles are dried and calcined. The drying is generally carried out at a temperature between room temperature up to 200° C, generally in the presence of air. The calcining is generally carried out at a temperature of 300 to 950° C, preferably 600 to 900° C, generally in the presence of air, for a period of 30 minutes to six hours. If so desired, the calcination may be carried out in the presence of steam to influence the crystal growth in the oxide.

By the above production method, it is possible to obtain a carrier having properties which will give a catalyst with the surface area, pore volume, and pore size distribution characteristics specified above. The surface area, pore volume, and pore size distribution characteristics can be adjusted in a manner know to the skilled person, for example by the addition during the mixing or shaping stage of an acid, such as nitric acid, acetic acid or formic acid, or other compounds as moulding auxiliary, or by regulating the water content of the gel by adding or removing water.

The carrier of the catalysts to be used in the process according to the invention have a specific surface area, pore volume, and pore size distribution of the same order as that of the catalysts themselves. The carrier of catalyst I preferably has a surface area of 100-200 m²/g, more preferably 130-170 m²/g. The total pore volume is preferably 0.5-1.2 ml/g, more preferably 0.7-1.0 ml/g. The carrier of catalyst II preferably has a surface area of 180-300 m²/g, more preferably 190-240 m²/g and a pore volume of 0.5-1.0 ml/g, more preferably 0.6-0.9 ml/g.

The Group VIIB metal components, Group VIII metal components, and, where appropriate, Group IA metal components, can be incorporated into the catalyst carrier in a conventional manner, e.g., by impregnation and/or by incorporation into the support material before it is shaped into particles. At this point in time it is considered preferred to first prepare the carrier and incorporate the catalytic materials into the carrier after it has been dried and calcined. The metal components can be incorporated into the catalyst composition in the form of suitable precursors, preferably by impregnating the catalyst with an acidic or basic impregnation solution comprising suitable metal precursors.

For the Group VIIB metals, ammonium heptamolybdate, ammonium dimolybdate, and ammonium tungstate may be mentioned as suitable precursors. Other compounds, such as oxides, hydroxides, carbonates, nitrates, chlorides, and organic acid salts, may also be used. For the Group VIII metals, suitable precursors include oxides, hydroxides, carbonates, nitrates, chlorides, and organic acid salts. Carbonates and nitrates are particularly suitable. Suitable Group IA metal precursors include nitrates and carbonates.

The impregnation solution, if applied, may contain other compounds the use of which is known in the art, such as organic acids, e.g., citric acid, ammonia water, hydrogen peroxide water, gluconic acid, tartaric acid, malic acid or EDTA (ethyleneediamine tetraacetic acid). It will be clear to the skilled person that there is a wide range of variations on this method. Thus, it is possible to apply a plurality of impregnating stages, the impregnating solutions to be used containing one or more of the component precursors that are to be deposited, or a portion thereof. Instead of impregnating techniques, dipping methods, spraying methods, etc. can be used. In the case of multiple impregnation, dipping, etc., drying and/or calcining may be carried out in between.

After the metals have been incorporated into the catalyst composition, it is optionally dried, e.g., in air flow for about 0.5 to 16 hours at a temperature between room temperature and 200° C, and subsequently calcined, generally in air, for about 1 to 6 hours, preferably 1-3 hours at 200-800° C, preferably 450-600° C. The drying is done to
physically remove the deposited water. The calcining is done to bring at least part, preferably all, of the metal component precursors to the oxide form.

[0057] It may be desirable to convert the catalyst, i.e., the Group VIII and Group VIII metal components present therein, into the sulfidic form prior to its use in the hydroprocessing of hydrocarbon feedstocks. This can be done in an otherwise conventional manner, e.g., by contacting the catalyst in the reactor at increasing temperature with hydrogen and a sulfur-containing feedstock, or with a mixture of hydrogen and hydrogen sulfide. Ex situ presulfiding is also possible.

[0058] The process of the present invention is particularly suitable for the hydropprocessing of heavy hydrocarbon feeds. It is particularly suitable for hydroprocessing of heavy feedstocks of which at least 50 wt. % boils above 538°C (1000°F) and which comprise at least 2 wt. % of sulfur and at least 5 wt. % of Conradson carbon. The sulfur content of the feedstock may be above 3 wt. %. Its Conradson carbon content may be above 8 wt. %. The feedstock may contain contaminants, such as nickel and vanadium. Typically, these metals are present in an amount of at least 20 wppm, calculated on the total of Ni and V, more particularly in an amount of at least 30 wppm.

[0059] Suitable feedstocks include atmospheric residue, vacuum residue, residues blended with gas oils, particularly vacuum gas oils, crude, shale oils, tar sand oils, solvent deasphalted oil, coal liquefied oil, etc. Typically they are atmospheric residue (AR), vacuum residue (VR), and mixtures thereof.

[0060] Each stage of the process according to the invention can be carried out in a fixed bed, in a moving bed, or in an ebullated bed. It is preferred to carry out at least one stage in an ebullating bed. Carrying out both stages in an ebullating bed is preferred.

[0061] The two stages of the process according to the invention can be carried out in a single reactor, or in separate reactors. If the process is in an ebullating bed process, the use of separate reactors for the two stages is preferred. If so desired one may perform one or more of intermediate phase separation, stripping, H₂ quenching, etc. between the two stages.

[0062] The process conditions for the process according to the invention may be as follows. The temperature generally is 350-450°C, preferably 400-440°C. The pressure generally is 5-25 MPa, preferably 14-19 MPa. The liquid hourly space velocity generally is 0.1-3 h⁻¹, preferably 0.3-2 h⁻¹. The hydrogen to feed ratio generally is 300-1,500 Nl/h, preferably 600-1,000 Nl/h. The process is carried out in the liquid phase.

[0063] The invention will be elucidated below by way of the following examples, though it must not be deemed limited thereby or thereto.

EXAMPLE 1

[0064] Preparation of Catalyst A

[0065] A sodium aluminate solution and an aluminum sulfate solution were simultaneously added dropwise to a tank containing tap water, mixed at pH 8.5 at 65°C, and held for 70 minutes. The thus produced alumina hydrate gel was separated from the solution and washed with warm water, to remove the impurities in the gel. Then, the gel was kneaded for about 20 minutes and extruded as cylindrical particles having a diameter of 0.9 to 1 mm and a length of 3.5 mm. The extruded alumina particles were dried at 120°C for 16 hours and calcined at 900°C for 2 hours, to obtain an alumina carrier.

[0066] 100 g of the alumina carrier obtained as described above were immersed in 100 ml of a citric acid solution containing 16.4 g of ammonium molybdate tetrahydrate and 9.8 g of nickel nitrate hexahydrate at 25°C for 45 minutes, to obtain a carrier loaded with metallic components.

[0067] Subsequently the loaded carrier was dried at 120°C for 30 minutes and calcined at 540°C for 1.5 hours, to complete a catalyst. The amounts of the respective components in the produced catalyst and the properties of the catalyst are shown in Table 1. Catalyst A meets the requirements of Catalyst I of the present invention.

[0068] Preparation of Catalyst B

[0069] To produce a silica-alumina carrier, a sodium aluminate solution was supplied to a tank containing tap water, and an aluminum sulfate solution was added and mixed. When the addition of the aluminum sulfate solution was completed, the mixture had a pH of 8.5. The mixture was kept at 64°C for 1.5 hours. Then, a sodium silicate solution was added and mixed. By such mixing a silica-alumina gel was produced. The sodium silicate concentration was set at 1.6 wt. % of the alumina gel solution.

[0070] The silica-alumina gel was isolated by filtration and washed with hot water to remove impurities from the gel. It was then extruded into cylindrical grains with a diameter of 0.9-1 mm and a length of 3.5 mm. The resulting particles were dried in air at a temperature of 120°C for 16 hours and subsequently calcined in the presence of air for two hours at 800°C to obtain a silica-alumina carrier. The silica-content of the obtained carrier was 7 wt. %.

[0071] One hundred grams of the thus obtained silica-alumina carrier were impregnated with 100 ml of an impregnation solution containing 13.4 g of ammonium molybdate tetrahydrate, 11.2 g of nickel nitrate hexahydrate, 0.68 g of sodium nitrate, and 50 ml of 25% ammonia water. The impregnated carrier was then dried at a temperature of 120°C for 30 minutes and calcined in a kiln for 1.5 hours at 540°C to produce a final catalyst. The composition and properties of this catalyst are given in Table 1. Catalyst B meets the requirements of Catalyst II of the present invention.

| TABLE 1 |
| --- | --- | --- |
| carrier | Catalyst A | Catalyst B |
| | alumina | Al₂O₃ + 6% SiO₂ |
| Group VIII wt. % ox | 11.9 | 11.5 |
| Group VIII wt. % ox | 2.0 | 2.1 |
| Group IX wt. % ox | 0 | 0.26 |
| surface area m²/g | 147 | 214 |
| total pore volume ml/g | 0.79 | 0.75 |
| % PV(≥200 Å) | 74 | 22.6 |
| % PV(100-200 Å) | 25 | 64 |
| % PV(100-1200 Å) | 89 | 75 |
| % PV(≥1000 Å) | 11 | 10 |
| % PV(≥5000 Å) | 0.4 | 12 |
Catalysts A and B were tested in various combinations in the two-stage hydroprocessing of a heavy hydrocarbon feedstock. The feedstock used in these examples was a Middle East petroleum consisting of 90 wt. % of vacuum residue (VR) and 10 wt. % of atmospheric residue (AR). The composition and properties of the feed are given in Table 1.

### TABLE 2

<table>
<thead>
<tr>
<th>Feedstock composition</th>
<th>Middle East petroleum (VR:AR = 50:50)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfur (wt. %)</td>
<td>4.79</td>
</tr>
<tr>
<td>Nitrogen (ppm)</td>
<td>2,890</td>
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<tr>
<td>Metals - vanadium (ppm)</td>
<td>85</td>
</tr>
<tr>
<td>Metals - nickel (ppm)</td>
<td>26</td>
</tr>
<tr>
<td>Conradson Carbon residue (wt. %)</td>
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<tr>
<td>C7-insolubles* (wt. %)</td>
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<td>Vacuum residue† (wt. %)</td>
<td>75</td>
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<tr>
<td>Density (g/ml at 15°C)</td>
<td>1.0048</td>
</tr>
</tbody>
</table>

*Matter insoluble in a-naphthalene

*Fraction boiling above 538°C in accordance with ASTM D 5307 (distillation gas chromatography)

Catalysts A and B were packed into a fixed bed reactor in two catalyst beds in the combinations given in Table 3 below. The two catalyst beds contained equal volume amounts of catalyst.

The cracking rate, also indicated as vacuum residuum cracking rate, is the conversion of the fraction boiling above 538°C into product boiling below 538°C.

<table>
<thead>
<tr>
<th>Catalyst I</th>
<th>C. 1</th>
<th>C. C. 1</th>
<th>C. C. 2</th>
<th>C. C. 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Catalyst II</td>
<td>B</td>
<td>A</td>
<td>B</td>
<td>B</td>
</tr>
<tr>
<td>HDS</td>
<td>115</td>
<td>100</td>
<td>126</td>
<td>116</td>
</tr>
<tr>
<td>HDM</td>
<td>112</td>
<td>100</td>
<td>76</td>
<td>85</td>
</tr>
<tr>
<td>Asphaltenes removal</td>
<td>107</td>
<td>100</td>
<td>81</td>
<td>82</td>
</tr>
</tbody>
</table>

In the above table it can be seen that, as compared with the comparative catalyst combinations, the catalyst system according to the invention combines good hydrodesulfurization, hydrometallization, and asphaltene removal with a high cracking rate and low sediment formation.

1. A method for hydrotreatment of a heavy hydrocarbon oil, comprising bringing a heavy hydrocarbon oil in a first stage into contact with hydrotreatment catalyst I in the presence of hydrogen, after which the effluent of the first stage is contacted in whole or in part with hydrotreatment catalyst II in the presence of hydrogen, wherein catalyst I comprises 7 to 20 wt. % of a Group VIB metal component, calculated as trioxide on the weight of the catalyst, and 0.5 to 6 wt. % of a Group VIII metal component, calculated as oxide on the weight of the catalyst, on a porous inorganic carrier, said catalyst having a specific surface area of at least 100 m²/g, a total pore volume of at least 0.55 ml/g, at least 50% of the total pore volume in pores with a diameter of at least 20 nm (200 Å) and at least 65% of the total pore volume in pores with a diameter of 10-120 nm (100-1200 Å), and catalyst II comprises 7 to 20 wt. % of a Group VIB metal component, calculated as trioxide on the weight of the catalyst, and 0.5 to 6 wt. % of a Group VIII metal component, calculated as oxide on the weight of the catalyst, on a porous inorganic carrier, said catalyst having a specific surface area of at least 100 m²/g, a total pore volume of at least 0.55 ml/g, 30-80% of the pore volume in pores with a diameter of 10-20 nm (100-200 Å), and at least 5% of the pore volume in pores with a diameter of at least 100 nm (1000 Å) with catalyst I having a larger percentage of its pore volume in pores with a diameter of at least 20 nm (200 Å) than catalyst II.

2. The method of claim 1 wherein catalyst I has less than 25% of its pore volume in pores with a diameter of less than 10 nm (100 Å).

3. The method of claim 1 wherein the carrier of catalyst I consists essentially of alumina.

4. The method of claim 1 wherein catalyst II has less than 50% of its pore volume in pores with a diameter of at least 20 nm (200 Å).

5. The method of claim 1 wherein the carrier of catalyst II consists essentially of alumina and at least 3.5 wt. % of silica.

6. The method of claim 1 wherein catalyst II comprises 0.1-2 wt. % of a Group IA metal component.

7. The method of claim 1 wherein the heavy hydrocarbon feed is a feed of which at least 50 wt. % boils above 538°C (1000°F) and which comprises at least 2 wt. % of sulfur and at least 5 wt. % of Conradson carbon.

8. The method of claim 1 wherein at least one of the stages is carried out in an ebullating bed.

9. A combination of catalysts comprising a catalyst I that comprises 7 to 20 wt. % of a Group VIB metal component, calculated as trioxide on the weight
of the catalyst, and 0.5 to 6 wt. % of a Group VIII metal component, calculated as oxide on the weight of the catalyst, on a porous inorganic carrier, said catalyst having a specific surface area of at least 100 m²/g, a total pore volume of at least 0.55 ml/g, at least 50% of the total pore volume in pores with a diameter of at least 20 nm (200 Å) and at least 65% of the total pore volume in pores with a diameter of 10-120 nm (100-1200 Å), and a catalyst II that comprises 7 to 20 wt. % of a Group VIIB metal component, calculated as trioxide on the weight of the catalyst, and 0.5 to 6 wt. % of a Group VIII metal component, calculated as oxide on the weight of the catalyst, on a porous inorganic carrier, said catalyst having a specific surface area of at least 100 m²/g, a total pore volume of at least 0.55 ml/g, 30-80% of the pore volume in pores with a diameter of 10-20 nm (100-200 Å), and at least 5% of the pore volume in pores with a diameter above 100 nm (1000 Å), with catalyst I having a larger percentage of its pore volume in pores with a diameter of at least 20 nm (200 Å) than catalyst II.

10. The catalyst combination of claim 9 wherein catalyst I has less than 25% of its pore volume in pores with a diameter of less than 10 nm (100 Å).

11. The catalyst combination of claim 9 wherein the carrier of catalyst I consists essentially of alumina.

12. The catalyst combination of claim 9 wherein catalyst II has less than 50% of its pore volume in pores with a diameter above 200 Å.

13. The catalyst combination of claim 9 wherein the carrier of catalyst II consists essentially of alumina and at least 3.5 wt. % of silica.

14. The catalyst combination of claim 9 wherein catalyst II comprises 0.1-2 wt. % of a Group IA metal component.

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