A two stage hydrodesulfurizing process for producing low sulfur distillates. A distillate boiling range feedstock containing in excess of about 3,000 ppm sulfur is hydrodesulfurized in a first hydrodesulfurizing stage containing one or more reaction zones in the presence of hydrogen and a hydrodesulfurizing catalyst. The liquid product stream thereof is passed to a first separation stage wherein a vapor phase product stream and a liquid product stream are produced. The liquid product stream, which has a substantially lower sulfur and nitrogen content than the original feedstream is passed to a second hydrodesulfurizing stage also containing one or more reaction zones where it is reacted in the presence of hydrogen and a second hydrodesulfurizing catalyst at hydrodesulfurizing conditions. The catalyst in any one or more reaction zones is a bulk multimetallic catalyst comprised of at least one Group VIII non-noble metal and at least two Group VIB metals.
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TWO STAGE PROCESS FOR HYDRODESULFURIZING DISTILLATES USING BULK MULTIMETALLIC CATALYST

CROSS REFERENCE TO RELATED APPLICATIONS

This is a continuation-in-part of USSN 09/231,156 filed on January 15, 1999, which is a continuation-in-part of USSN 08/900,389 which was filed on July 15, 1997.

FIELD OF THE INVENTION

This invention relates to a two stage hydrodesulfurizing process for producing low sulfur distillates. A distillate boiling range feedstock containing in excess of about 3,000 wppm sulfur is hydrodesulfurized in a first hydrodesulfurizing stage containing one or more reaction zones in the presence of hydrogen and a hydrodesulfurizing catalyst. The liquid product stream thereof is passed to a first separation stage wherein a vapor phase product stream and a liquid product stream are produced. The liquid product stream, which has a substantially lower sulfur and nitrogen content then the original feedstream is passed to a second hydrodesulfurizing stage also containing one or more reaction zones where it is reacted in the presence of hydrogen and a second hydrodesulfurizing catalyst at hydrodesulfurizing conditions. The catalyst in any one or more reaction zones is a bulk multimetallic catalyst comprised of at least one Group VIII non-noble metal and at least two GroupVIB metals wherein the ratio of Group VIB metal to Group VIII non-noble metals is about 10:1 to 1:10.

BACKGROUND OF THE INVENTION

Environmental and regulatory initiatives are requiring ever lower levels of both sulfur and aromatics in distillate fuels. For example, proposed sulfur limits for distillate fuels to be marketed in the European Union for the year 2005 is 50
wppm or less. There are also regulations that will require lower levels of total aromatics in hydrocarbons and, more specifically, to lower levels of multiring aromatics found in distillate fuels and heavier hydrocarbon products. Further, the maximum allowable aromatics level for U.S. on-road diesel, CARB reference diesel, and Swedish Class I diesel are 35, 10 and 5 vol.%, respectively. Further, the CARB and Swedish Class I diesel fuels allow no more than 1.4 and 0.02 vol.% polyaromatics, respectively. Consequently, much work is presently being done in the hydrotreating art because of these proposed regulations.

Hydrotreating, or in the case of sulfur removal, hydrodesulfurization, is well known in the art and typically requires treating the petroleum streams with hydrogen in the presence of a supported catalyst at hydrotreating conditions. The catalyst is usually comprised of a Group VI metal with one or more Group VIII metals as promoters on a refractory support. Hydrotreating catalysts that are particularly suitable for hydrodesulfurization, as well as hydrodenitrogenation, generally contain molybdenum or tungsten on alumina promoted with a metal such as cobalt, nickel, iron, or a combination thereof. Cobalt promoted molybdenum on alumina catalysts are most widely used when the limiting specifications are hydrodesulfurization, while nickel promoted molybdenum on alumina catalysts are the most widely used for hydrodenitrogenation, partial aromatic saturation, as well as hydrodesulfurization.

Much work is also being done to develop more active catalysts and improved reaction vessel designs in order to meet the demand for more effective hydroprocessing processes. Various improved hardware configurations have been suggested. One such configuration is a countercurrent design wherein the feedstock flows downwardly through successive catalyst beds counter to upflowing treat gas, which is typically a hydrogen containing treat-gas. The downstream catalyst beds, relative to the flow of feed, can contain high performance, but otherwise more sulfur sensitive catalysts because the
upflowing treat gas carries away heteroatom components. such as H₂S and NH₃. that are deleterious to sulfur and nitrogen sensitive catalysts.

One approach to prepare improved hydrotreating catalysts is, a family of compounds. related to hydrotalcites. e.g., ammonium nickel molybdates, has been prepared. Whereas X-ray diffraction analysis has shown that hydrotalcites are composed of layered phases with positively charged sheets and exchangeable anions located in the galleries between the sheets, the related ammonium nickel molybdate phase has molybdate anions in interlayer galleries bonded to nickel oxyhydroxide sheets. See, for example, Levin, D., Soled, S. L., and Ying, J. Y.. Crystal Structure of an Ammonium Nickel Molybdate prepared by Chemical Precipitation. Inorganic Chemistry, Vol. 35, No. 14, p. 4191-4197 (1996). The preparation of such materials also has been reported by Teichner and Astier. Appl. Catal. 72, 321-29 (1991); Ann. Chim. Fr. 12, 337-43 (1987). and C. R. Acad. Sci. 304 (II), #11, 563-6 (1987) and Mazzocchia, Solid State Ionics, 63-65 (1993) 731-35.

Now, when molybdenum is partially substituted for by tungsten, an amorphous phase is produced which upon decomposition and, preferably, sulfidation, provides enhanced hydrodenitrogenation (HDN) catalyst activity relative to the unsubstituted (Ni-Mo) phase.

SUMMARY OF THE INVENTION

In accordance with this invention there is provided a two stage hydrodesulfurizing process for reducing the sulfur content of distillate feedstocks having a sulfur content greater than about 3.000 wppm. which process comprises:
a) reacting said feedstream in a first hydodesulfurization stage in the presence of a hydrogen-containing treat gas. said first hydrotreating stage containing one or more reaction zones, each reaction zone operated at hydodesulfurizing conditions and in the presence of a hydodesulfurization catalyst, thereby resulting in a liquid product stream having a sulfur content less than about 3,000 wppm;

b) passing the liquid product stream of said first hydodesulfurization stage to a separation zone where a vapor phase product stream and a liquid phase product stream are produced;

c) reacting said liquid phase product stream of b) above in a second hydodesulfurization stage in the presence of a hydrogen-containing treat gas said second hydodesulfurization stage containing one or more reaction zones operated at hydodesulfurization conditions wherein each reaction zone contains a bed of hydrotreating catalyst, thereby resulting in a liquid product stream having less than about 1,000 wppm sulfur;

d) passing the liquid product stream of step c) above to a separation zone wherein a vapor phase stream and a liquid phase stream are produced;

e) collecting both said vapor phase stream and said liquid phase stream; and

wherein at least one of the reaction zones of at least one of said hydodesulfurizing stages contains a bulk multimetallic catalyst comprised of at least one Group VIII non-noble metal and at least two Group VIB metals and wherein the ratio of Group VIB metal to Group VIII non-noble metal is from about 10:1 to about 1:10.
In a preferred embodiment of the present invention the Group VIII non-noble metal is selected from Ni and Co and the Group VIB metals are selected from Mo and W.

In another preferred embodiment of the present invention two Group VIB metals are present as Mo and W and the ratio of Mo to W is about 9:1 to about 1:9.

In yet another preferred embodiment of the present invention the bulk multimetallic is represented by the formula:

\[(X)_b \ (Mo)_c \ (W)_d \ O_z\]

wherein X is a Group VIII non-noble metal, and the molar ratio of b: (c+d) is 0.5/1 to 3/1, preferably 0.75/1 to 1.5/1, more preferably 0.75/1 to 1.25/1.

In still another preferred embodiment of the present invention the molar ratio of c:d is preferably >0.01/1, more preferably >0.1/1, still more preferably 1/10 to 10/1, still more preferably 1/3 to 3/1, most preferably substantially equimolar amounts of Mo and W, e.g., 2/3 to 3/2; and z = \([2b + 6 \ (c+d)]/2\).

In another preferred embodiment of the present invention the essentially amorphous material has a unique X-ray diffraction pattern showing crystalline peaks at d = 2.53 Angstroms and d = 1.70 Angstroms.

In still another preferred embodiment of the present invention the Group VIII non-noble metal is nickel.

In a preferred embodiment of the present invention the bulk multimetallic catalyst also contains an acid function

In still another preferred embodiment of the present invention at least a portion of the vapor phase product stream from the first separation zone is recycled to the first hydrodesulfurization stage.
In another preferred embodiment of the present invention at least a portion of the vapor phase product stream from the second separation stage is cascaded to said first hydrodesulfurization stage.

**BRIEF DESCRIPTION OF DRAWINGS**

Figure 1 is the X-ray diffraction pattern of a NH₄-Ni₁.₅Mo₀.₅W₀.₅ compound prepared by boiling precipitation before calcining (Curve A) and after calcining at 400°C (Curve B). Note that the patterns for both the precursor and the decomposition product of the precursor are quite similar with the two peaks at essentially the same place. The ordinate is relative intensity; the abscissa is two theta (degrees).

Figure 2 shows the X-ray diffraction patterns, by CuKα radiation (λ=1.5405Å), of NH₄-Ni-Moₓₜₓ-Wₓ-O precursors wherein curve A is Mo₀.₉W₀.₁, curve B is Mo₀.₇W₀.₃, curve C is Mo₀.₅W₀.₅, curve D is Mo₀.₃W₀.₇, curve E is Mo₀.₁W₀.₉, and curve F is Mo OW₁. The ordinate and abscissa are as described for Figure 1.

**DETAILED DESCRIPTION OF THE INVENTION**

The bulk multimetallic catalyst composition used in the practice of the present invention can be used in virtually all hydroprocessing processes to treat a plurality of feeds under wide-ranging reaction conditions such as temperatures of from 200 to 450°C, hydrogen pressures of from 5 to 300 bar, liquid hourly space velocities of from 0.05 to 10 h⁻¹ and hydrogen treat gas rates of from 35.6 to 1780 m³/m³ (200 to 10000 SCF/B). The term "hydroprocessing" encompasses all processes in which a hydrocarbon feed is reacted with hydrogen at the temperatures and pressures noted above, and include hydrodemetallation, hydrodewaxing, hydrotreating, hydrogenation hydrodesulfurization.
hydrodenitrogenation, hydrodearomatization, hydroisomerization, and hydrocracking including selective hydrocracking. Depending on the type of hydroprocessing and the reaction conditions, the products of hydroprocessing may show improved viscosities, viscosity indices, saturates content, low temperature properties, volatilities and depolarization. It is to be understood that hydroprocessing of the present invention can be practiced in one or more reaction zones and can be practiced in either countercurrent flow or cocurrent flow mode. By countercurrent flow mode we mean a process mode wherein the feedstream flows countercurrent to the flow of hydrogen-containing treat gas. The hydroprocessing reactor can also be operated in any suitable catalyst-bed arrangement mode. For example, it can be a fixed bed, slurry bed, or ebulating bed.

A wide range of petroleum and chemical feedstocks can be hydroprocessed in accordance with the present invention. Suitable feedstocks include whole and reduced petroleum crudes, atmospheric and vacuum residua, asphaltenes, deasphalted oils, cycle oils, FCC tower bottoms, gas oils, including atmospheric and vacuum gas oils and coker gas oils, light to heavy distillates including raw virgin distillates, hydrocrackates, hydrotreated oils, dewaxed oils, slack waxes, Fischer-Tropsch waxes, raffinates, naphthas, and mixtures thereof.

The instant invention can be practiced by feeding a distillate boiling range feedstock containing high levels of sulfur and nitrogen to a first hydrodesulfurization reaction stage to remove a substantial amount of the sulfur and nitrogen. Suitable feeds are those containing in excess of about 3,000 wppm sulfur and are typically raw virgin distillates. The feed is hydrodesulfurized in a first stage, which contains one or more reaction zones, in the presence of hydrogen and a first hydrotreating catalyst (described below) of the present invention under hydrodesulfurizing conditions. The product stream is passed to
a separation zone wherein a vapor phase stream and a liquid phase stream are produced. The liquid phase product stream is passed to a second hydrodesulfurization stage, which also contains one or more reaction zones, where it is further hydrodesulfurized in the presence of hydrogen and a second hydrodesulfurization catalyst. The liquid product stream from the second hydrodesulfurization stage is passed to a second separation zone wherein a vapor product stream and liquid product stream are produced and collected for further processing or blending. It is within the scope of this invention that at least a portion of the vapor product stream from either or both reaction stages can be recycled to the first or second reaction stage. The vapor product, which contains hydrogen treat gas can also be cascaded from the second reaction stage to the first reaction stage.

The feedstock to the first hydrodesulfurization stage will typically contain greater than about 3,000 wppm sulfur. The product stream from the first stage hydrodesulfurization will typically have a sulfur content of about 300 to about 1,500 wppm, preferably from about 300 to 1,000 wppm, and more preferably from about 300 to 750 wppm. The product stream from the second stage hydrodesulfurization will have a sulfur content less than about 150 wppm, preferably less than about 100 wppm, and more preferably less than about 50 wppm. As previously mentioned, one or more of the reaction zones in one or both of the hydrodesulfurization stages will contain the bulk multimetallic catalyst of the invention. The second hydrodesulfurization stage must contain, in at least one reaction zone, a bed of the bulk multimetallic catalyst of the present invention. The first hydrodesulfurization stage may or may not contain a bed of the bulk catalyst of this invention. The first hydrodesulfurization stage can only conventional hydrodesulfurization catalyst or it can contain one reaction zone with conventional hydrodesulfurization catalyst and at least one reaction zone that contains the bulk multimetallic catalyst of the present invention. For example, one or both of the reactors of the reaction stages can
contain two or more reaction zones, each reaction zone containing catalyst. The first reaction stage can contain a bed of conventional hydrodesulfurization catalyst in one reaction zone and a bed of the bulk multimetallic catalyst of this invention in another reaction zone. Preferably the first reaction zone will contain the conventional hydrodesulfurization catalyst. The second hydrodesulfurization stage in addition to the bulk multimetallic catalyst of the present invention can contain a bed of isom-enhanced bulk multimetallic catalyst. It is also with the scope of this invention that the bulk multimetallic catalyst of this invention can be used in all reaction zones of both reaction stages.

The hydrotreating catalyst of any one or more of the reaction zones of any one or both of the hydrodesulfurization stages is a bulk multimetallic catalyst comprised of at least one Group VIII non-noble metal and at least two Group VIB metals and wherein the ratio of Group VIB metal to Group VIII non-noble metal is from about 10:1 to about 1:10. It is preferred that the catalyst be a bulk trimetallic catalyst comprised of one Group VIII non-noble metal, preferably Ni or Co and the two Group VIB metals Mo and W. It is preferred that the ratio of Mo to W be about 9:1 to about 1:9.

The preferred bulk trimetallic catalyst compositions used in the practice of the present invention is represented by the formula:

\[(X)_b (Mo)_c (W)_d O_z\]

wherein X is one or more Group VIII non-noble metal, the molar ratio of b: (c+d) is 0.5/1 to 3/1, preferably 0.75/1 to 1.5/1, more preferably 0.75/1 to 1.25/1;

The molar ratio of c:d is preferably >0.01/1. more preferably >0.1/1, still more preferably 1/10 to 10/1, still more preferably 1/3 to 3/1. most preferably
substantially equimolar amounts of Mo and W. e.g., 2/3 to 3/2; and \( z = \frac{2b + 6 (c+d)}{2} \).

The essentially amorphous material has a unique X-ray diffraction pattern showing crystalline peaks at \( d = 2.53 \) Angstroms and \( d = 1.70 \) Angstroms.

The mixed metal oxide is readily produced by the decomposition of a precursor having the formula:

\[
(NH_3)_a (X)_{2b} (Mo)_{c} (W)_d O_z
\]

wherein the molar ratio of \( a:b \) is \( \leq 1.0/1 \), preferably \( 0-1 \); and \( b, c, \) and \( d \), are as defined above, and \( z = \frac{a + 2b + 6 (c+d)}{2} \). The precursor has similar peaks at \( d = 2.53 \) and \( 1.70 \) Angstroms.

Decomposition of the precursor may be effected at elevated temperatures, e.g., temperatures of at least about 300°C, preferably about 300-450°C, in a suitable atmosphere, e.g., inerts such as nitrogen, argon, or steam, until decomposition is substantially complete, i.e., the ammonium is substantially completely driven off. Substantially complete decomposition can be readily established by thermogravimetric analysis (TGA), i.e., flattening of the weight change curve.

The catalyst compositions used in the practice of the present invention can be prepared by any suitable means. One such means is a method wherein not all of the metals are in solution. Generally, the contacting of the metal components in the presence of the protic liquid comprises mixing the metal component and subsequently reacting the resulting mixture. It is essential to the solid route that at least one metal component is added at least partly in the solid state during the mixing step and that the metal of at least one of the metal
components which have been added at least partly in the solid state. remains at least partly in the solid state during the mixing and reaction step. "Metal" in this context does not mean the metal in its metallic form but present in a metal compound, such as the metal component as initially applied or as present in the bulk catalyst composition.

Generally, during the mixing step either at least one metal component is added at least partly in the solid state and at least one metal component is added in the solute state, or all metal components are added at least partly in the solid state, wherein at least one of the metals of the metal components which are added at least partly in the solid state remains at least partly in the solid state during the entire process of the solid route. That a metal component is added "in the solute state" means that the whole amount of this metal component is added as a solution of this metal component in the protic liquid. That a metal component is added "at least partly in the solid state" means that at least part of the metal component is added as solid metal component and, optionally, another part of the metal component is added as a solution of this metal component in the protic liquid. A typical example is a suspension of a metal component in a protic liquid in which the metal is at least partly present as a solid, and optionally partly dissolved in the protic liquid.

To obtain a bulk catalyst composition with high catalytic activity, it is therefore preferred that the metal components, which are at least partly in the solid state during contacting, are porous metal components. It is desired that the total pore volume and pore size distribution of these metal components is approximately the same as those of conventional hydrotreating catalysts. Conventional hydrotreating catalysts generally have a pore volume of 0.05 - 5 ml/g, preferably of 0.1 - 4 ml/g, more preferably of 0.1 - 3 ml/g and most preferably of 0.1 - 2 ml/g determined by nitrogen adsorption. Pores with a
diameter smaller than 1 nm are generally not present in conventional hydrotreating catalysts. Further, conventional hydrotreating catalysts have generally a surface area of at least 10 m²/g and more preferably of at least 50 m²/g and most preferably of at least 100 m²/g, determined via the B.E.T. method. For instance, nickel carbonate can be chosen which has a total pore volume of 0.19 - 0.39 ml/g and preferably of 0.24 - 0.35 ml/g determined by nitrogen adsorption and a surface area of 150 - 400 m²/g and more preferably of 200 - 370 m²/g determined by the B.E.T. method. Furthermore these metal components should have a median particle diameter of at least 50 nm, more preferably at least 100 nm, and preferably not more than 5000 μm and more preferably not more than 3000 μm. Even more preferably, the median particle diameter lies in the range of 0.1 - 50 μm and most preferably in the range of 0.5 - 50 μm. For instance, by choosing a metal component which is added at least partly in the solid state and which has a large median particle diameter, the other metal components will only react with the outer layer of the large metal component particle. In this case, so-called "core-shell" structured bulk catalyst particles are obtained.

An appropriate morphology and texture of the metal component can either be achieved by applying suitable preformed metal components or by preparing these metal components by the above-described precipitation under such conditions that a suitable morphology and texture is obtained. A proper selection of appropriate precipitation conditions can be made by routine experimentation.

As has been set out above, to retain the morphology and texture of the metal components which are added at least partly in the solid state, it is essential that the metal of the metal component at least partly remains in the solid state during the whole process of this solid route. It is noted again that it is essential
that in no case should the amount of solid metals during the process of the solid route becomes zero. The presence of solid metal comprising particles can easily be detected by visual inspection at least if the diameter of the solid particles in which the metals are comprised is larger than the wavelength of visible light. Of course, methods such as quasi-elastic light scattering (QELS) or near forward scattering which are known to the skilled person can also be used to ensure that in no point in time of the process of the solid route, all metals are in the solute state.

The protic-liquid to be applied in the solid or solution route of this invention for preparing catalyst can be any protic liquid. Examples include water, carboxylic acids, and alcohols such as methanol or ethanol. Preferably, a liquid comprising water such as mixtures of an alcohol and water and more preferably water is used as protic liquid in this solid route. Also different protic liquids can be applied simultaneously in the solid route. For instance, it is possible to add a suspension of a metal component in ethanol to an aqueous solution of another metal component.

The Group VIB metal generally comprises chromium, molybdenum, tungsten, or mixtures thereof. Suitable Group VIII non-noble metals are, e.g., iron, cobalt, nickel, or mixtures thereof. Preferably, a combination of metal components comprising nickel, molybdenum and tungsten or nickel, cobalt, molybdenum and tungsten is applied in the process of the solid route. If the protic liquid is water, suitable nickel components which are at least partly in the solid state during contacting comprise water-insoluble nickel components such as nickel carbonate, nickel hydroxide, nickel phosphate, nickel phosphite, nickel formate, nickel sulfide, nickel molybdate, nickel tungstate, nickel oxide, nickel alloys such as nickel-molybdenum alloys, Raney nickel, or mixtures thereof. Suitable molybdenum components, which are at least partly in the solid state
during contacting, comprise water-insoluble molybdenum components such as molybdenum (di- and tri) oxide, molybdenum carbide, molybdenum nitride, aluminum molybdate, molybdic acid (e.g. H₂MoO₄), molybdenum sulfide, or mixtures thereof. Finally, suitable tungsten components which are at least partly in the solid state during contacting comprise tungsten di- and trioxide, tungsten sulfide (WS₂ and WS₃), tungsten carbide, tungstic acid (e.g. H₂WO₄ - H₂O, H₂W₄O₁₃ - 9H₂O), tungsten nitride, aluminum tungstate (also meta-, or polytungstate) or mixtures thereof. These components are generally commercially available or can be prepared by, e.g., precipitation. e.g., nickel carbonate can be prepared from a nickel chloride. sulfate, or nitrate solution by adding an appropriate amount of sodium carbonate. It is generally known to the skilled person to choose the precipitation conditions in such a way as to obtain the desired morphology and texture.

In general, metal components, which mainly contain C, O. and/or H beside the metal, are preferred because they are less detrimental to the environment. Nickel carbonate is a preferred metal component to be added at least partly in the solid state because when nickel carbonate is applied, CO₂ evolves and positively influences the pH of the reaction mixture. Further, due to the transformation of carbonate into CO₂, the carbonate does not end up in the wastewater.

Preferred nickel components which are added in the solute state are water-soluble nickel components, e.g. nickel nitrate, nickel sulfate, nickel acetate, nickel chloride, or mixtures thereof. Preferred molybdenum and tungsten components which are added in the solute state are water-soluble molybdenum and tungsten components such as alkali metal or ammonium molybdate (also peroxo-, di-, tri-, tetra-, hepta-, octa-, or tetradecamolybdate), Mo-P heteropolyanion compounds, Wo-Si heteropolyanion compounds, W-P
heteropolyanion compounds, W-Si heteropolyanion compounds, Ni-Mo-W heteropolyanion compounds, Co-Mo-W heteropolyanion compounds, alkali metal or ammonium tungstates (also meta-, para-, hexa-, or polytungstate), or mixtures thereof.

Preferred combinations of metal components are nickel carbonate, tungstic acid and molybdenum oxide. Another preferred combination is nickel carbonate, ammonium dimolybdate and ammonium metatungstate. It is within the scope of the skilled person to select further suitable combinations of metal components. It must be noted that nickel carbonate always comprises a certain amount of hydroxy-groups. It is preferred that the amount of hydroxy-groups present in the nickel carbonate be high.

An alternative method of preparing the catalysts used in the practice of the present invention is to prepare the bulk catalyst composition by a process comprising reacting in a reaction mixture a Group VIII non-noble metal component in solution and a Group VIB metal component in solution to obtain a precipitate. As in the case of the solid route, preferably, one Group VIII non-noble metal component is reacted with two Group VIB metal components. The molar ratio of Group VIB metals to Group VIII non-noble metals applied in the process of the solution route is preferably the same as described for the solid route. Suitable Group VIB and Group VIII non-noble metal components are, e.g. those water-soluble nickel, molybdenum and tungsten components described above for the solid route. Further Group VIII non-noble metal components are, e.g., cobalt or iron components. Further Group VIB metal components are, e.g. chromium components. The metal components can be added to the reaction mixture in solution, suspension or as such. If soluble salts are added as such, they will dissolve in the reaction mixture and subsequently be precipitated. Suitable Group VIB metal salts which are soluble in water are ammonium salts
such as ammonium dimolybdate, ammonium tri-, tetra- hepta-, octa-, and tetradeca- molybdate, ammonium para-, meta-, hexa-, and polytungstate, alkali metal salts, silicic acid salts of Group VIB metals such as molybdic silicic acid, molybdc silicic tungstic acid, tungstic acid, metatungstic acid, pertungstic acid, heteropolyanion compounds of Mo-P, Mo-Si, W-P, and W-Si. It is also possible to add Group VIB metal-containing compounds which are not in solution at the time of addition, but where solution is effected in the reaction mixture. Examples of these compounds are metal compounds which contain so much crystal water that upon temperature increase they will dissolve in their own metal water. Further, non-soluble metal salts may be added in suspension or as such, and solution is effected in the reaction mixture. Suitable non-soluble metals salts are heteropolyanion compounds of Co-Mo-W (moderately soluble in cold water), heteropolyanion compounds of Ni-Mo-W (moderately soluble in cold water).

The reaction mixture is reacted to obtain a precipitate. Precipitation is effected by adding a Group VIII non-noble metal salt solution at a temperature and pH at which the Group VIII non-noble metal and the Group VIB metal precipitate, adding a compound which complexes the metals and releases the metals for precipitation upon temperature increase or pH change or adding a Group VIB metal salt solution at a temperature and pH at which the Group VIII non-noble metal and Group VIB metal precipitate, changing the temperature, changing the pH, or lowering the amount of the solvent. The precipitate obtained with this process appears to have high catalytic activity. In contrast to the conventional hydroprocessing catalysts, which usually comprise a carrier impregnated with Group VIII non-noble metals and Group VIB metals, said precipitate can be used without a support. Unsupported catalyst compositions are usually referred to as bulk catalysts. Changing the pH can be done by adding base or acid to the reaction mixture, or adding compounds, which decompose upon temperature, increase into hydroxide ions or H⁺ ions that respectively
increase or decrease the pH. Examples of compounds that decompose upon
temperature increase and thereby increase or decrease the pH are urca, nitrites,
ammonium cyanate, ammonium hydroxide, and ammonium carbonate.

In an illustrative process according to the solution route, solutions of
ammonium salts of a Group VIB metal are made and a solution of a Group VIII
non-noble metal nitrate is made. Both solutions are heated to a temperature of
approximately 90°C. Ammonium hydroxide is added to the Group VIB metal
solution. The Group VIII non-noble metal solution is added to the Group VIB
metal solution and direct precipitation of the Group VIB and Group VIII
non-noble metal components occurs. This process can also be conducted at
lower temperature and/or decreased pressure or higher temperature and/or
increased pressure.

In another illustrative process according to the solution route, a Group
VIB metal salt, a Group VIII metal salt, and ammonium hydroxide are mixed in
solution together and heated so that ammonia is driven off and the pH is lowered
to a pH at which precipitation occurs. For instance when nickel, molybdenum,
and tungsten components are applied, precipitation typically occurs at a pH
below 7.

Independently from whether the solid or solution route is chosen in step
(i), the resulting bulk catalyst composition preferably comprises and more
preferably consists essentially of bulk catalyst particles having the characteristics
of the bulk catalyst particles described under the heading "Catalyst compositions
of the invention."

The bulk catalyst composition can generally be directly shaped into
hydroprocessing particles. If the amount of liquid of the bulk catalyst
composition is so high that it cannot be directly subjected to a shaping step. A solid liquid separation can be performed before shaping. Optionally the bulk catalyst composition, either as such or after solid liquid separation, can be calcined before shaping.

The median diameter of the bulk catalyst particles is at least 50 nm, more preferably at least 100 nm, and preferably not more than 5000 μm and more preferably not more than 3000 μm. Even more preferably, the median particle diameter lies in the range of 0.1 - 50 μm and most preferably in the range of 0.5-50μm.

If a binder material is used in the preparation of the catalyst composition, it can be any material that is conventionally applied as a binder in hydroprocessing catalysts. Examples include silica, silica-alumina, such as conventional silica-alumina, silica-coated alumina and alumina-coated silica, alumina such as (pseudo)boehmite, or gibbsite, titania, zirconia, cationic clays or anionic clays such as saponite, bentonite, kaoline, sepiolite or hydrotalcite, or mixtures thereof. Preferred binders are silica, silica-alumina, alumina, titanic, zirconia, or mixtures thereof. These binders may be applied as such or after peptization. It is also possible to apply precursors of these binders that, during the process of the invention are converted into any of the above-described binders. Suitable precursors are, e.g., alkali metal aluminates (to obtain an alumina binder), water glass (to obtain a silica binder), a mixture of alkali metal aluminates and water glass (to obtain a silica alumina binder), a mixture of sources of a di-, tri-, and/or tetravalent metal such as a mixture of water-soluble salts of magnesium, aluminum and/or silicon (to prepare a cationic clay and/or anionic clay), chlorohydrol, aluminum sulfate, or mixtures thereof.

If desired, the binder material may be composited with a Group VIB metal and/or a Group VIII non-noble metal, prior to being composited with the
bulk catalyst composition and/or prior to being added during the preparation thereof. Compositing the binder material with any of these metals may be carried out by impregnation of the solid binder with these materials. The person skilled in the art knows suitable impregnation techniques. If the binder is peptized, it is also possible to carry out the peptization in the presence of Group VIIB and/or Group VII non-noble metal components.

If alumina is applied as binder, the surface area preferably lies in the range of 100 - 400 m²/g, and more preferably 150 - 350 m²/g, measured by the B.E.T. method. The pore volume of the alumina is preferably in the range of 0.5 - 1.5 ml/g measured by nitrogen adsorption.

Generally, the binder material to be added in the process of the invention has less catalytic activity than the bulk catalyst composition or no catalytic activity at all. Consequently, by adding a binder material, the activity of the bulk catalyst composition may be reduced. Therefore, the amount of binder material to be added in the process of the invention generally depends on the desired activity of the final catalyst composition. Binder amounts from 0 - 95 wt.% of the total composition can be suitable, depending on the envisaged catalytic application. However, to take advantage of the resulting unusual high activity of the composition of the present invention, binder amounts to be added are generally in the range of 0.5 - 75 wt.% of the total composition.

The catalyst composition can be directly shaped. Shaping comprises extrusion, pelleting, beading, and/or spray drying. It must be noted that if the catalyst composition is to be applied in slurry type reactors, fluidized beds, moving beds, expanded beds, or ebulliating beds, spray drying or beading is generally applied for fixed bed applications. Generally, the catalyst composition is extruded, pelleted and/or beaded. In the latter case, prior to or during the
shaping step, any additives that are conventionally used to facilitate shaping can be added. These additives may comprise aluminum stearate, surfactants, graphite or mixtures thereof. These additives can be added at any stage prior to the shaping step. Further, when alumina is used as a binder, it may be desirable to add acids prior to the shaping step such as nitric acid to increase the mechanical strength of the extrudates.

It is preferred that a binder material is added prior to the shaping step. Further, it is preferred that the shaping step is carried out in the presence of a liquid, such as water. Preferably, the amount of liquid in the extrusion mixture, expressed as LOI is in the range of 20 - 80%.

The resulting shaped catalyst composition can, after an optional drying step, be optionally calcined. Calcination however is not essential to the process of the invention. If a calcination is carried out in the process of the invention, it can be done at a temperature of, e.g., from 100° - 600°C and preferably 350° to 500°C for a time varying from 0.5 to 48 hours. The drying of the shaped particles is generally carried out at temperatures above 100°C.

In a preferred embodiment of the invention, the catalyst composition is subjected to spray drying, (flash) drying, milling, kneading, or combinations thereof prior to shaping. These additional process steps can be conducted either before or after a binder is added, after solid-liquid separation, before or after calcination and subsequent to re-wetting. It is believed that by applying any of the above-described techniques of spray drying, (flash) drying, milling, kneading, or combinations thereof, the degree of mixing between the bulk catalyst composition and the binder material is improved. This applies to both cases where the binder material is added before or after the application of any of the above-described methods. However, it is generally preferred to add the
binder material prior to spray drying and/or any alternative technique. If the binder is added subsequent to spray drying and/or any alternative technique, the resulting composition is preferably thoroughly mixed by any conventional technique prior to shaping. An advantage of, e.g., spray drying is that no wastewater streams are obtained when this technique is applied.

Furthermore, a cracking component may be added during catalyst preparation. The cracking component may serve as an isomerization enhancer. The cracking component can be any conventional cracking component such as cationic clays, anionic clays, zeolites such as ZSM-5, (ultra-stable) zeolite Y, zeolite X, ALPO's, SAPO's, amorphous cracking components such as silica-alumina, or mixtures thereof. It will be clear that some materials may act as a binder and a cracking component at the same time. For instance, silica-alumina may have at the same time a cracking and a binding function.

If desired, the cracking component may be composited with a Group VIB metal and/or a Group VIII non-noble metal prior to being composited with the bulk catalyst composition and/or prior to being added during the preparation thereof. Compositing the cracking component with any of these metals may be carried out by impregnation of the cracking component with these materials.

The cracking component, which can comprise about 0-80 wt.%, based on the total weight of the catalyst, can be added at any stage of the process of the present invention prior to the shaping step. However, it is preferred to add the cracking component during the compositing step (ii) with the binder.

It is within the scope of the present invention is the use of the catalysts of this invention admixed with a Pt/Pd containing zeolite for restoring octane number during hydrodesulfurization on cat naphthas. The instant invention can also be used to reduce the acid number of petroleum crudes.
Generally, it depends on the envisaged catalytic application of the final catalyst composition which of the above-described cracking components is added. A zeolite is preferably added if the resulting composition shall be applied in hydrocracking or fluid catalytic cracking. Other cracking components such as silica-alumina or cationic clays are preferably added if the final catalyst composition shall be used in hydrotreating applications. The amount of cracking material that is added depends on the desired activity of the final composition and the application envisaged and thus may vary from 0 - 80 wt.%, based on the total weight of the catalyst composition.

If desired, further materials can be added in addition to the metal components already added. These materials include any material that is added during conventional hydroprecipitation catalyst preparation. Suitable examples are phosphorus compounds, boron compounds, fluorine-containing compounds, additional transition metals, rare earth metals, fillers, or mixtures thereof.

Suitable phosphorus compounds include ammonium phosphate, phosphoric acid, or organic phosphorus compounds. Phosphorus compounds can be added at any stage of the process of the present invention prior to the shaping step and/or subsequent to the shaping step. If the binder material is peptized, phosphorus compounds can also be used for peptization. For instance, the binder can be peptized by contacting the binder with phosphoric acid or with a mixture of phosphoric and nitric acid.

Suitable additional transition metals are, e.g., rhenium, ruthenium, rhodium, iridium, chromium, vanadium, iron, cobalt, platinum, palladium, cobalt, nickel, molybdenum, or tungsten. Nickel, molybdenum, and tungsten can be applied in the form of any of the water-insoluble nickel, molybdenum and/or tungsten components that are described above for the solid route. These
metals can be added at any stage of the process of the present invention prior to the shaping step. Apart from adding these metals during the process of the invention, it is also possible to composite the final catalyst composition therewith. It is, e.g., possible to impregnate the final catalyst composition with an impregnation solution comprising any of these metals.

The processes of the present invention for preparing the bulk catalyst compositions may further comprise a sulfidation step. Sulfidation is generally carried out by contacting the catalyst composition or precursors thereof with a sulfur containing compound such as elementary sulfur, hydrogen sulfide or polysulfides. The sulfidation can generally be carried out subsequently to the preparation of the bulk catalyst composition but prior to the addition of a binder material, and/or subsequently to the addition of the binder material but prior to subjecting the catalyst composition to spray drying and/or any alternative method, and/or subsequently to subjecting the composition to spray drying and/or any alternative method but prior to shaping, and/or subsequently to shaping the catalyst composition. It is preferred that the sulfidation is not carried out prior to any process step that reverts the obtained metal sulfides into their oxides. Such process steps are, e.g., calcination or spray drying or any other high temperature treatment in the presence of oxygen. Consequently, if the catalyst composition is subjected to spray drying and/or any alternative technique, the sulfidation should be carried out subsequent to the application of any of these methods.

Additionally to, or instead of, a sulfidation step, the bulk catalyst composition may be prepared from at least one metal sulfide. If, e.g. the solid route is applied in step (i), the bulk catalyst component can be prepared from nickel sulfide and/or molybdenum sulfide and/or tungsten sulfide.
If the catalyst composition is used in a fixed bed processes, the sulfidation is preferably carried out subsequent to the shaping step and, if applied, subsequent to the last calcination step. Preferably, the sulfidation is carried out \textit{ex situ}, i.e., the sulfidation is carried out in a separate reactor prior to loading the sulfided catalyst composition into the hydroprocessing unit. Furthermore, it is preferred that the catalyst composition is both sulfided \textit{ex situ} and \textit{in situ}.

One or more of the reaction zones of any or both of the hydrodesulfurization stages, may contain a conventional hydrodesulfurization catalyst. Suitable conventional hydrodesulfurization catalysts for use in the present invention includes those that are comprised of at least one Group VIII metal, preferably Fe, Co or Ni, more preferably Co and/or Ni, and most preferably Co; and at least one Group VI metal, preferably Mo or W, more preferably Mo, on a relatively high surface area support material, preferably alumina. Other suitable hydrodesulfurization catalyst supports include zeolites, amorphous silica-alumina, and titania-alumina. Noble metal catalysts can also be employed, preferably when the noble metal is selected from Pd and Pt. It is within the scope of the present invention that more than one type of hydrodesulfurization catalyst be used in the same reaction vessel. The Group VIII metal is typically present in an amount ranging from about 2 to 20 wt.%, preferably from about 4 to 12%. The Group VI metal will typically be present in an amount ranging from about 5 to 50 wt.%, preferably from about 10 to 40 wt.%, and more preferably from about 20 to 30 wt.%. All metals weight percents are on support. By "on support" we mean that the percents are based on the weight of the support. For example, if the support were to weigh 100 g, then 20 wt.% Group VIII metal would mean that 20 g of Group VIII metal was on the support.

It has been found that in this case, the bulk catalyst particles are sintering-resistant. Thus the active surface area of the bulk catalyst particles is
maintained during use. The molar ratio of Group VIB to Group VIII non-noble metals ranges generally from 10:1 - 1:10 and preferably from 3:1 - 1:3. In the case of a core-shell structured particle, these ratios of course apply to the metals contained in the shell. If more than one Group VIB metal is contained in the bulk catalyst particles, the ratio of the different Group VIB metals is generally not critical. The same holds when more than one Group VIII non-noble metal is applied. In the case where molybdenum and tungsten are present as Group VIB metals, the molybdenum:tungsten ratio preferably lies in the range of 9:1 - 1:9. Preferably the Group VIII non-noble metal comprises nickel and/or cobalt. It is further preferred that the Group VIB metal comprises a combination of molybdenum and tungsten. Preferably, combinations of nickel/molybdenum/tungsten and cobalt/molybdenum/tungsten and nickel/cobalt/molybdenum/tungsten are used. These types of precipitates appear to be sinter-resistant. Thus, the active surface area of the precipitate is remained during use.

The metals are preferably present as oxidic compounds of the corresponding metals, or if the catalyst composition has been sulfided, sulfidic compounds of the corresponding metals.

Preferably the particles have a surface area of at least 50 m²/g and more preferably of at least 100 m²/g measured VIB the B.E.T. method. It is furthermore preferred that the particles comprise 50 - 100 wt.%, and even more preferably 70 - 100 wt.% of at least one Group VIII non-noble metal and at least one Group VIB metal, based on the total weight of the particles, calculated as metal oxides. The amount of Group VIB and Group VIII non-noble metals can easily be determined VIB TEM-EDX.

It is desired that the pore size distribution of the particles is approximately the same as the one of conventional hydrotreating catalysts. More in particular, these
particles have preferably a pore volume of 0.05 - 5 ml/g, more preferably of 0.1 - 4 ml/g, still more preferably of 0.1 - 3 ml/g and most preferably 0.1 - 2 ml/g determined by nitrogen adsorption. Preferably, pores smaller than 1 nm are not present. Furthermore these particles preferably have a median diameter of at least 50 nm, more preferably at least 100 nm, and preferably not more than 5000 μm and more preferably not more than 3000 μm. Even more preferably, the median particle diameter lies in the range of 0.1 - 50 μm and most preferably in the range of 0.5 - 50 μm.

The surface area of the catalyst composition preferably is at least 40 m²/g, more preferably at least 80 m²/g and most preferably at least 120 m²/g. The total pore volume of the catalyst composition is preferably at least 0.05 ml/g and more preferably at least 0.01 ml/g as determined by water porosimetry. To obtain catalyst compositions with high mechanical strength, it may be desirable that the catalyst composition of the invention has a low macroporosity.

It was found that the bulk catalyst particles have a characteristic X-ray diffraction pattern which differs from catalysts obtained by co-mixing and conventional hydroprocessing catalysts obtained by impregnation. The X-ray diffraction pattern of the bulk catalyst particles comprises, and preferably essentially consists of, peaks characteristic to the reacted metal components. If, e.g., nickel hydroxy-carbonate has been contacted with a molybdenum and tungsten component as described above, the resulting bulk catalyst particles are characterized by an X-ray diffraction pattern which comprises peaks at d values of (4.09), 2.83, 2.54, 2.32, 2.23, 1.71, (1.54), 1.47. Values in brackets indicate that the corresponding peaks are rather broad and/or have a low intensity or are not distinguished at all. The term "the X-ray diffraction pattern essentially consists of " these peaks means that apart from these peaks, there are essentially no further peaks contained in the diffraction pattern. The precipitate for catalyst obtained by the solution route has a characteristic X-ray diffraction pattern
which differs from catalyst obtained by co-mixing and conventional hydroprocessing catalysts obtained by impregnation. For instance the X-ray diffraction pattern of a Ni-Mo-W precipitate as prepared by the solution route has peaks at d values of 2.52, 1.72 and 1.46.

Also as previously stated, the catalyst composition may comprise conventional hydroprocessing catalysts. The binder materials and cracking components of the conventional hydroprocessing catalyst generally comprise any of the above-described binder materials and cracking components. The hydrogenation metals of the conventional hydroprocessing catalyst generally comprise Group VIB and Group VIII non-noble metals such as combinations of nickel or cobalt with molybdenum or tungsten. Suitable conventional hydroprocessing catalysts are, e.g., hydrotreating catalysts. These catalysts can be in the spent, regenerated, or fresh state.

As will be clear from the above, it is possible to add the Group VIII non-noble metal containing compound and the Group VIB metal-containing compound in various ways, at various temperatures and pHs, in solution, in suspension, and as such, simultaneously and sequentially.

The precursor compound can also be readily prepared by one of several methods, including a variation of the boiling decomposition method used by Teichner and Astier in which a tungsten compound is added to the initial mixture of a molybdenum salt, a nickel salt and ammonium hydroxide. Direct precipitation and pH controlled precipitation may also be used to prepare the precursor compound. In all cases, however, water soluble salts of nickel, molybdenum and tungsten are employed.
Preferably, the molybdenum and tungsten salts are ammonium compounds, e.g., ammonium molybdate, ammonium metatungstate, while the nickel salt may be the nitrate or hydrated nitrates.

The decomposed precursor can be sulfided or pre-sulfided by a variety of known methods. For example, the decomposition product can be contacted with a gas comprising H₂S and hydrogen, e.g., 10% H₂S/H₂, at elevated temperatures for a period of time sufficient to sulfide the decomposition product, usually at the point of H₂S breakthrough in the exit gas. Sulfiding can also be effected in situ, by passing a typical feedstock containing sulfur over the decomposition product.

Process conditions applicable for the use of the catalysts described herein may vary widely depending on the feedstock to be treated. Thus, as the boiling point of the feed increases, the severity of the conditions will also increase. The following table serves to illustrate typical conditions for a range of feeds.

<table>
<thead>
<tr>
<th>FEED</th>
<th>TYPICAL BOILING RANGE °C</th>
<th>TEMP. °C</th>
<th>PRESS. BAR</th>
<th>SPACE VELOCITY V/V/HR</th>
<th>H₂ GAS RATE SCF/B</th>
</tr>
</thead>
<tbody>
<tr>
<td>naphtha</td>
<td>25-210</td>
<td>100-370</td>
<td>10-60</td>
<td>0.5-10</td>
<td>100-2,000</td>
</tr>
<tr>
<td>diesel</td>
<td>170-350</td>
<td>200-400</td>
<td>15-110</td>
<td>0.5-4</td>
<td>500-6,000</td>
</tr>
<tr>
<td>heavy gas oil</td>
<td>325-475</td>
<td>260-430</td>
<td>15-170</td>
<td>0.3-2</td>
<td>1000-6,000</td>
</tr>
<tr>
<td>lube oil</td>
<td>290-550</td>
<td>200-450</td>
<td>6-210</td>
<td>0.2-5</td>
<td>100-10,000</td>
</tr>
<tr>
<td>residuum</td>
<td>10-50%&gt;575</td>
<td>340-450</td>
<td>65-1100</td>
<td>0.1-1</td>
<td>2,000-10,000</td>
</tr>
</tbody>
</table>

The high activity bulk multimetallic HDN and isom-enhanced HDS catalysts of the present invention could be optimized for use in a countercurrent flow reactor. A countercurrent flow reactor offers several advantages over co-
current reactors including: high hydrogen partial pressures; internal stripping of 
H₂S and NH₃; and high reactor temperature. One non-limiting process option 
employing countercurrent flow technology is to feed to raw feedstock in a first 
hydrodesulfurization stage which is operated in co-current mode to produce a 
product which is lower in sulfur and nitrogen than the original feed. The 
hydrotreated oil is then flashed to separate light gases, especially ammonia, from 
the liquid product. If necessary, the hydrocarbon portion of this interstage vapor 
stream can be hydrotreated in a dedicated “vapor-phase” reactor or condensed 
and reintroduced into the first-stage reactor to reduce sulfur to acceptable levels. 
The hydrotreated feed is then processed over an isom-enhanced HDS catalyst in 
a countercurrent reactor. Liquid feed is introduced at the top of the reactor while 
fresh hydrogen is added at the bottom. Product ammonia and hydrogen sulfide 
are stripped by treat gas and exit the top of the reactor. Some of the liquid feed 
will vaporize and pass overhead. This vapor stream can be hydrotreated in a 
“vapor phase” bed above the feed injection point in the countercurrent flow 
reactor or condensed and reintroduced into the first-stage reactor to reduce sulfur 
to acceptable levels.

While the two stage hydrodesulfurization process of the present invention 
is applicable to all sulfur-containing compounds common to petroleum and 
chemical streams, the isom-enhanced catalyst of the process is particularly 
useful for the conversion of the least reactive, most highly refractory sulfur 
species, especially the class derived from dibenzothiophenes, and most 
especially the alkyl-substituted derivatives of this heterocyclic group.

The following examples will serve to illustrate, but not limit, this invention.

**Example 1** Preparation of NH₄-Ni-Mo-O Phase (boiling decomposition as per 
Teichner and Astier procedure):
In a 1 liter flask, 26.5 g ammonium molybdate (0.15 moles Mo) and 43.6 g nickel nitrate hexahydrate (0.15 moles Ni) were dissolved in 300 cc of water so that the resulting pH equaled 4.3. To this solution, a concentrated NH₄OH solution was added. At first, a precipitate formed which on further addition of NH₄OH dissolved to give a clear blue solution with a pH of 8.3, and additional NH₄OH (~250cc) was added until a pH of 10 was reached. The solution was heated to 90°C for 3 h during which ammonia gas evolved and a green precipitate formed. The final pH lay between 6.8 and 7. The suspension was cooled to room temperature, filtered, washed with water and dried at 120°C overnight. About 18.6g of material was obtained. The sample analyzed for Ni at 26.6 wt.% and Mo at 34 wt.%. The X-ray diffraction spectra of the phase matches the pattern reported by Teichner.

**Example 2.** Preparation of NH₄-Ni-Mo₅W₅-O by boiling decomposition:

In a 1 liter flask, 13.2 g ammonium molybdate (0.075 moles Mo), 18.7 g ammonium metatungstate (0.075 moles W) and 43.6 g nickel nitrate hexahydrate (0.15 moles Ni) were dissolved in 300cc of water so that the resulting pH equaled 4.3. To this solution, a concentrated NH₄OH solution (~600cc) was added until the pH reached 10. At this point, some precipitate remained. The solution was refluxed at ~100°C for 3 h. During this heating, the precipitate dissolved to give a clear blue solution and on further heating, a green precipitate formed. The heating was continued until the pH reached between 6.8 and 7. The suspension was cooled to room temperature, filtered, washed with water and dried at 120°C overnight. 18 grams of material is obtained. The X-ray diffraction spectra of the phase is given in Figure 2 showing an amorphous background with the two largest peaks at d=2.58 and 1.70Å.
**Example 3** Preparation of NH$_4$-Ni-Mo$_{0.5}$W$_{0.5}$-O by direct precipitation:

In a 1 liter flask, 17.65 g of ammonium molybdate (0.1 mole Mo) and 24.60 g of ammonium metatungstate (0.1 mole W) were dissolved in 800 cc of water giving a solution pH of ~5.2. To this solution 0.4 moles of NH$_4$OH (~30 cc) was added, raising the pH to ~9.8 (solution A). This solution was warmed to 90°C. A second solution was prepared by adding 58.2 g of nickel nitrate, (0.2 moles Ni) which was dissolved in 50 cc of water (solution B) and maintained at 90°C. This solution was added dropwise at a rate of 7 cc/min into the ammonium molybdate/ammonium metatungstate solution. A precipitate begins to form after 1/4 of the solution was added. This suspension which was at a pH ~6.5 was stirred for 30 minutes while the temperature was maintained at 90°C. The material was filtered hot, washed with hot water, and dried at 120°C. Approximately 38 g of material was recovered.

**Example 4** Preparation of NH$_4$-Ni-Mo$_{0.5}$-Mo$_{0.5}$W$_{0.5}$-O by controlled pH precipitation:

Two solutions were prepared with the same amounts of nickel, tungsten, molybdenum and ammonium hydroxide are described in Example 3 (solutions A and B) except that each solution contained about 700 cc of water. The two solutions were added into a separate vessel initially containing 400 cc of water held at 90°C. Solution B (the acidic solution) was pumped into the vessel at a constant rate of ~15cc/min, while solution A is added through a separate pump which is under feedback PC control and set to maintain the pH at 6.5. On mixing the two solutions a precipitate forms. The slurry was stirred at 90°C for 30 minutes, filtered hot, washed with hot water, and dried at 120°C.

**Example 5** Catalytic Evaluation Using Dibenzothiophene (DBT):
1.5-2 g of the catalysts of Examples 1-4 were placed in a quartz boat which was in turn inserted into a horizontal quartz tube and placed into a Lindberg furnace. The temperature was raised to 370°C in about one hour with N₂ flowing at 50 cc/m, and the flow continued for 1.5 h at 370°C. N₂ was switched off and 10% H₂S/H₂ then added to the reactor at 20 cc/m. the temperature increased to 400°C, and held there for 2 hours. The heat was then shut off and the catalyst cooled in flowing H₂S/H₂ to 70°C, at which point this flow was discontinued and N₂ was added. At room temperature, the quartz tube was removed and the material transferred into a N₂ purged glove box. Catalysts were evaluated in a 300cc modified Carberry batch reactor designed for constant hydrogen flow. The catalyst was pilled and sized to 20/40 mesh and one gram was loaded into a stainless steel basket, sandwiched between a layer of mullite beads. 100 cc of liquid feed containing 5 wt.% dibenzothiophene in decalin was added to the autoclave. A hydrogen flow of 100 cc/min was passed through the reactor and the pressure was maintained at 3150kPa using a back pressure regulator. The temperature was raised to 350°C at 5-6 deg/min and run until either 50% DBT was converted or until 7 hours was reached. A small aliquot of product was removed every 30 minutes and analyzed by GC. Rate constants for the overall conversion as well as the conversion to the reaction products biphenyl (BP) and cyclohexylbenzene (CHB) were calculated as described by M. Daage and R. R. Chianelli [J. Cat. 149, 414-27 (1994)] and are shown in Table 1. As described in that article, high selectivities to cyclohexylbenzene relative to BP during the desulfurization reaction are a good indication of a catalyst with high hydrodenitrogenation activity, whereas high selectivities of BP relative to CHB indicates a catalyst with high hydrodesulfurization activity.

The results show that partial substitution of tungsten for molybdenum results in catalysts that are substantially higher for DBT conversion. A standard
supported Ni-Mo on Al₂O₃ catalyst is also shown for comparison. The high CHB/BP ratio suggests that the catalysts are active for HDN.

Table 1. Comparison of Activity in DBT Conversion Tests With Tungsten Addition by Different Preparation Schemes

<table>
<thead>
<tr>
<th>catalyst</th>
<th>preparation technique</th>
<th>example #</th>
<th>K_{total} @ 350°C</th>
<th>CHB/BP @ 350°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH₄-Ni-Mo-O</td>
<td>boiling decomposition</td>
<td>1</td>
<td>106</td>
<td>10.4</td>
</tr>
<tr>
<td>NH₄-Ni-Mo₅W₅-O</td>
<td>boiling decomposition</td>
<td>2</td>
<td>171</td>
<td>10.2</td>
</tr>
<tr>
<td>NH₄-Ni-Mo₅W₅-O</td>
<td>direct precipitation</td>
<td>3</td>
<td>167</td>
<td>12.4</td>
</tr>
<tr>
<td>NH₄-Ni-Mo₅W₅-O</td>
<td>controlled pH preparation</td>
<td>4</td>
<td>181</td>
<td>12.0</td>
</tr>
<tr>
<td>NiMo/Al₂O₃</td>
<td>impregnation</td>
<td></td>
<td>129</td>
<td>6.4</td>
</tr>
</tbody>
</table>

Example 6

A series of catalysts were prepared in accordance with the general preparation scheme of example 2 (i.e., boiling decomposition) but varying the Mo and W relative ratios by changing the amount of ammonium molybdate and ammonium metatungstate added to the solutions. Decomposition was effected as described in Example 5. The catalysts so prepared are shown in Table 2 along with their catalytic activities for DBT measured as described in Example 5.

Table 2. Comparison of Activity in DBT Conversion Tests with Variation in Relative W and Mo content

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Sample</th>
<th>ammonium molybdate (g)</th>
<th>ammonium metatungstate (g)</th>
<th>nickel nitrate hexahydrate (g)</th>
<th>K_{total} @ 350°C</th>
<th>CHB/BP @ 350°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH₄-NiW-O</td>
<td>18983-97</td>
<td>0</td>
<td>36.95</td>
<td>43.62</td>
<td>128</td>
<td>11.3</td>
</tr>
<tr>
<td>NH₄-NiMo₄W₉-O</td>
<td>18983-125</td>
<td>2.65</td>
<td>33.62</td>
<td>43.62</td>
<td>132</td>
<td>14.1</td>
</tr>
<tr>
<td>NH₄-NiMo₅W₇-O</td>
<td>18983-101</td>
<td>7.94</td>
<td>25.87</td>
<td>43.62</td>
<td>154</td>
<td>11.6</td>
</tr>
</tbody>
</table>
NH₄-NiMo₀.₅W₀.₅-O  18357-109  13.17  18.74  43.62  171  10.2
NH₄-NiMo₀.₇W₀.₃-O  18983-95  18.54  11.09  43.62  158  11.5
NH₄-NiMo₀.₉W₀.₁-O  18983-92  23.83  3.69  43.62  141  10.5

The data show that the most active catalyst contains an approximately equimolar mixture of tungsten and molybdenum.

**Example 7**

A series of catalysts were prepared as described in Example 3 (direct precipitation) in which equimolar mixtures of Mo and W were precipitated but the nickel content was varied. Decomposition was effected as described in Example 5. The catalysts so prepared are shown in Table 3 along with their catalytic activities for DBT measured as described in example 5.

**Table 3.** Variation of Nickel Content in NH₄-Ni-Mo₀.₅W₀.₅-O Catalysts

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Sample</th>
<th>ammonium molybdate (g)</th>
<th>ammonium metatungstate (g)</th>
<th>nickel nitrate hexahydrate (g)</th>
<th>K_total @ 350°C</th>
<th>CHB/B P @ 350°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH₄-Ni₀.₇5Mo₀.₅W₀.₅-O</td>
<td>19086-110</td>
<td>17.65</td>
<td>24.6</td>
<td>43.65</td>
<td>171</td>
<td>13.0</td>
</tr>
<tr>
<td>NH₄-Ni₁.₀Mo₀.₅W₀.₅-O</td>
<td>19086-82</td>
<td>17.65</td>
<td>24.6</td>
<td>58.2</td>
<td>167</td>
<td>12.4</td>
</tr>
<tr>
<td>NH₄-Ni₁.₂₅Mo₀.₅W₀.₅-O</td>
<td>19086-111</td>
<td>17.65</td>
<td>24.6</td>
<td>72.75</td>
<td>174</td>
<td>11.0</td>
</tr>
<tr>
<td>NH₄-Ni₁.₅Mo₀.₅W₀.₅-O</td>
<td>19086-112</td>
<td>17.65</td>
<td>24.6</td>
<td>87.3</td>
<td>148</td>
<td>9.55</td>
</tr>
</tbody>
</table>

Catalytic performance does not change substantially with variations in Ni from 0.75 to 1.5, although K appears to go through a maximum at about 1.25 Ni.

**Example 8**
A series of catalysts were prepared in which the quantity of NH₄OH used in the preparation was varied. The catalysts were prepared in accordance to the procedure described in Example 3 except that the amount of NH₄OH in solution A was varied to change to NH₄OH/Ni molar ratio when the two solutions were mixed. Decomposition was effected as described in Example 5. The catalysts so prepared are shown in Table 4 along with their catalytic activities for DBT measured as described in Example 5.

Table 4. Variation in NH₄OH Addition to Preparation

<table>
<thead>
<tr>
<th>Catalyst NH₄OH /Ni molar ratio</th>
<th>Sample</th>
<th>ammonium molybdate (g)</th>
<th>ammonium metatungstate (g)</th>
<th>nickel nitrate hexahydrate (g)</th>
<th>cm³ conc NH₄OH @ 350ºC</th>
<th>K_total @ 350ºC</th>
<th>K_CHB/ BP @ 350ºC</th>
</tr>
</thead>
<tbody>
<tr>
<td>1:2</td>
<td>19086-96</td>
<td>17.65</td>
<td>24.6</td>
<td>43.65</td>
<td>6.8</td>
<td>102</td>
<td>10.5</td>
</tr>
<tr>
<td>1:1</td>
<td>19086-97</td>
<td>17.65</td>
<td>24.6</td>
<td>58.2</td>
<td>14</td>
<td>137</td>
<td>10.4</td>
</tr>
<tr>
<td>2:1</td>
<td>19086-82</td>
<td>17.65</td>
<td>24.6</td>
<td>72.75</td>
<td>30</td>
<td>167</td>
<td>12.4</td>
</tr>
<tr>
<td>3:1</td>
<td>19086-104</td>
<td>17.65</td>
<td>24.6</td>
<td>87.3</td>
<td>41</td>
<td>164</td>
<td>11.4</td>
</tr>
<tr>
<td>4:1</td>
<td>19086-106</td>
<td>17.65</td>
<td>24.6</td>
<td>87.3</td>
<td>.55</td>
<td>161</td>
<td>12.1</td>
</tr>
</tbody>
</table>

While decomposition of the precursor compound will drive off most, if not all, of the ammonium portion of the precursor, the preparation of the precursor and the catalytic utility of the decomposition product can be affected by the amount of NH₄OH employed. Thus, the effectiveness of the decomposition product as a catalyst is enhanced when the NH₄OH/Ni ratio in the preparation of the precursor compound is from about 1:1 to about 4:1, preferably about 1.5:1 to about 4:1, and more preferably about 2:1 to about 4:1. While not wishing to be bound by any particular theory or mechanism, there is some evidence the NH₄OH/Ni ratio causes the Ni-M-W-O phase to change in the decomposition product.
Example 9

The catalysts of examples 1 and 2 were compared against standard supported Ni-Mo catalysts for the conversion of a LSADO (low sulfur auto diesel oil feed). This feed contained 510 wppm sulfur, 50 wppm nitrogen, and 30.6% aromatics with a gravity of 39.8° API. The catalysts were tested at 579°F, 650 psig of H₂, and 1850 SCFB/B of H₂. The relative activities of the different catalysts are summarized in Table 5.

Table 5. Relative Hydrotreating Activities on LSADO Feed

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Relative Volumetric HDS Activity</th>
<th>Relative Volumetric HDN Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni₅Mo/Al₂O₃</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>NH₄-NiMo-O</td>
<td>0.25</td>
<td>0.50</td>
</tr>
<tr>
<td>NH₄-Ni0.5Mo₅W₅-O</td>
<td>1.4</td>
<td>2.05</td>
</tr>
</tbody>
</table>

The Ni₅Mo/Al₂O₃ catalyst is a standard HDN/HDS catalyst, the NH₄-Ni-Mo phase is the bulk phase with no tungsten, and the NH₄-Ni₁.₀Mo₅W₅-O is the bulk phase with partial substitution of W for Mo. The NH₄-NiMo-O catalyst is also representative of known compounds. The catalyst of this invention is illustrated by NH₄-Ni₁.₀Mo₀.₅W₀.₅-O and the data show the clear advantage of ammonium nickel tungsten molybdate for HDN activity.

Example 10

Preparation of a bulk catalyst composition according to the solid route:
18.1kg-ammonium dimolybdate (15.33kg MoO₃) are dissolved in 575 liters water. Subsequently 28.5kg ammonium metatungstate (24 69kg WO₃) is added to the solution. The resulting solution is preheated up to 90°C. 26.5kg NiCO₃
(49.7% Ni) powder is mixed with water and the resulting paste is added to the ammonium dimolybdate/ammonium metatungstate solution. The resulting mixture is reacted for 7 hours at 89°C.

Example 11

Preparation of a bulk catalyst composition according to the solution route:
In a 1-liter flask, 13.2 g ammonium molybdate (0.075 moles Mo), 18.7 g ammonium metatungstate (0.075 moles W) and 43.6 g nickel nitrate hexahydrate (0.15 moles Ni) were dissolved in 300 ml water so that the resulting pH equaled 4.3. To this solution, a concentrated NH₄OH solution (about 600 ml) was added until the pH reached 10. At this point, some precipitate remained. The solution was refluxed at 100°C for 3 hours. During this heating, the precipitate dissolved to give a clear blue solution and on further heating, a green precipitate formed. The heating was continued until the pH reached a value between 6.8 and 7.0. The suspension was cooled to room temperature, filtered, washed with water and dried at 120°C overnight. 18 grams of material were obtained.

Example 12 (sample 2110587)

657g of a NiMo-W bulk catalyst composition obtained according to the procedure described in Examples 10 or 11 was added to 1362 g of an aqueous slurry containing 125g of alumina (prepared by precipitation of sodium aluminate and aluminum sulfate). The resulting Ni-Mo-W bulk catalyst - alumina composition was subsequently mixed at 80°C until an LOI of 31% was obtained. The resulting composition was subsequently extruded and the extrudates were dried at 120°C for about 90 minutes and subsequently calcined at 385°C for one hour in air.

Example 13 (sample 2110598)
The process of Example 12 was repeated except mat instead of the alumina suspension, a silica sol containing 10 wt.% silica were applied.

**Example 14 (sample 2110591)**

657g of a Ni-Mo-W bulk catalyst composition obtained according to the procedure described in Examples 7 or 8 was added to 510g of a boehmite paste containing 125g boehmite. The rebuffing paste was mixed at 60°C to obtain an LOI of 42%. The resulting composition was extruded, dried and calcined as described in Example 12.

**Example 15 (sample 2110469).**

The procedure described in Example 7 or 8 was repeated except that alumina is present during the preparation of the bulk catalyst composition. To 755g of the resulting dried Ni-Mo-W bulk catalyst - alumina composition containing 60g alumina, 461g water and a small amount of nitric acid were added. The resulting mixture was mixed at 70°C while evaporating water until an LOI of 34% was obtained. The resulting composition was extruded, dried and calcined as described in Example 12.

**Example 16**

Ammonium molybdate, ammonium tungsten and/or ammonium chromate are dissolved and combined in a first reactor. The temperature is increased to 90°C. The Group VIII salt is dissolved in a second reactor and heated to 90°C. Ammonium hydroxide is added to the first reactor to form a basic solution. The Group VIII metal solution is added to the first dropwise with stirring in 20 minutes. After 30 minutes, the precipitate is filtered and washed. The precipitate is dried overnight at 120°C and calcined at 385°C.
**Example 17**

The precipitation method of Example 16 was used to prepare a precipitate from ammonium dimolybdate, ammonium meta tungstate and Fe(III)(NO₃)₃ • 9 H₂O in 98% yield comprising 41.2 wt.% Fe₂O₃, 21.3 wt.% MoO₃, and 36.9 wt.% WO₃. The surface area of the precipitate was 76 m²/g. The pore volume as measured up to 60 nm by BET using the adsorption curve was 0.147 ml/g.

**Example 18**

The precipitation method of Example 16 was used to prepare a precipitate from Ni(CO₃)₂ • 6H₂O, (NH₄)₆Mo₇O₂₄•4H₂O, and (NH₄)₂Cr₂O₇ in 87.7% yield comprising 52.2 wt.% NiO, 29.4 wt.% MoO₃, and 16.6 wt.% Cr₂O₃. The surface area of the precipitate was 199 m²/g. The pore volume as measured up to 60 nm by BET using the adsorption curve was 0.276 ml/g.

**Example 19**

The precipitation method of Example 16 was used to prepare a precipitate from Ni(CO₃)₂ • 6H₂O, (NH₄)₆H₂W₁₂O₄₀, and (NH₄)₂Cr₂O₇ in 87.7% yield comprising 44.0 wt.% NiO, 42.4 wt.% WO₃, and 11.8 wt.% Cr₂O₃. The surface area of the precipitate was 199 m²/g. The pore volume as measured up to 60 nm by BET using the adsorption curve was 0.245 ml/g.

**Example 20 (High HDN Activity)**

A reactor was charged with 4.0 cc of 14/35 mesh bulk multimetallic catalyst of the present invention designated BMCat and having a composition Ni₁₃Mo₀.₅W₀.₅. After liquid phase sulfiding, the catalyst was used to process a hydrotreated virgin distillate with API gravity of 36.0 containing 510 wppm sulfur and 61 wppm nitrogen. Results are presented in Table 6.
Example 21 (Comparative)

A reactor was charged with a 4.0 cc of 14/35 mesh convention CoMo on alumina catalyst commercially available from Akzo Nobel as KF756. After liquid phase sulfiding, the catalyst was used to process the same hydrotreated virgin distillate as in Example 20. Results are presented in Table 6.

### Table 6

Comparison of HDS and HDN Activities for KF756 and BMCat

<table>
<thead>
<tr>
<th>Example</th>
<th>Catalyst</th>
<th>Temperature (°C)</th>
<th>Pressure (psig)</th>
<th>Product S (wppm)</th>
<th>RVA* For HDS</th>
<th>Product N (wppm)</th>
<th>RVA* For HDN</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
<td>RT-721#</td>
<td>282</td>
<td>650</td>
<td>165</td>
<td>217</td>
<td>4</td>
<td>223</td>
</tr>
<tr>
<td>2a</td>
<td>KF756#</td>
<td>282</td>
<td>650</td>
<td>280</td>
<td>--</td>
<td>18</td>
<td>--</td>
</tr>
<tr>
<td>1b</td>
<td>RT-721</td>
<td>282</td>
<td>400</td>
<td>230</td>
<td>180</td>
<td>9</td>
<td>283</td>
</tr>
<tr>
<td>2b</td>
<td>KF756</td>
<td>282</td>
<td>400</td>
<td>315</td>
<td>--</td>
<td>31</td>
<td>--</td>
</tr>
<tr>
<td>1c</td>
<td>RT-721</td>
<td>304</td>
<td>400</td>
<td>80</td>
<td>194</td>
<td>6</td>
<td>272</td>
</tr>
<tr>
<td>2c</td>
<td>KF756</td>
<td>304</td>
<td>400</td>
<td>160</td>
<td>--</td>
<td>26</td>
<td>--</td>
</tr>
<tr>
<td>1d</td>
<td>RT-721</td>
<td>325</td>
<td>400</td>
<td>20</td>
<td>211</td>
<td>4</td>
<td>279</td>
</tr>
<tr>
<td>2d</td>
<td>KF756</td>
<td>325</td>
<td>400</td>
<td>60</td>
<td>--</td>
<td>23</td>
<td>--</td>
</tr>
<tr>
<td>1e</td>
<td>RT-721#</td>
<td>282</td>
<td>650</td>
<td>165</td>
<td>203</td>
<td>4</td>
<td>517</td>
</tr>
<tr>
<td>2e</td>
<td>KF756#</td>
<td>282</td>
<td>650</td>
<td>270</td>
<td>--</td>
<td>36</td>
<td>--</td>
</tr>
</tbody>
</table>

1.0 LHSV and 1900 SCF/B H₂

*Relative to KF756 (1.5 order HDS and 1.0 order HDN rate basis)

# Start-of-run

End-of-run

HDN rate enhancement equal to about 280 RVA was demonstrated at 400 psig. Rates for HDS were also higher, with about a 200 RVA compared to KF756. The high HDN rates enable the use of a nitrogen-sensitive, ultra-high activity HDS catalysts in the second-stage. Results obtained on a nitrogen-free feed are described below.

Example 22 (High HDS Activity)

The hydrotreated distillate of Example 20 was subjected to further hydrotreating to produce a feed with a nil sulfur and nitrogen, and an API gravity of 36.9. This was spiked to 460 wppm sulfur with 4,6-diethyl-debenzothiophene. A reactor
charged with an intimate mixture of 3.76 g (4.8 cc) of 14/35 mesh KF756 and 0.37 g (1.0 cc) of 0.9 Pt/ECR-32(13) zeolite was used to process the distillate feed described above. Results are presented in Table 7.

**Example 23 (High HDS Activity)**

A reactor charged with an intimate mixture of 3.71 g (4.8cc) of 14/35 mesh KF756 and 0.82 g (2.0 cc) of 0.9 Pt/ECR-32(13) zeolite (19487-5A) was used to process the feed of Example 22. Results are presented in Table 7.

**Example 24 (Comparative)**

A reactor was charged with 3.96 g (5.1cc) of 14/35 mesh KF756 and was used to process the feed of Example 22. Results are presented in Table 7.

**Table 7**

*Comparison of HDS for KF756 and KF756 + Solid Acid*

<table>
<thead>
<tr>
<th>Example</th>
<th>Catalyst(s)</th>
<th>Product S (wppm)</th>
<th>RVA* For HDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>8:1 (w/w) KF756 + Pt/ECR-32(13)</td>
<td>90</td>
<td>260</td>
</tr>
<tr>
<td>4</td>
<td>4:1 (w/w) KF756 + Pt/ECR-32(13)</td>
<td>30</td>
<td>371</td>
</tr>
<tr>
<td>5</td>
<td>KF756</td>
<td>265</td>
<td>--</td>
</tr>
</tbody>
</table>

315°C, 300 psig, 2.0 LHSV (over KF756) and 1000 SCF/B H₂

*Relative to KF756 alone (1.0 order HDS rate basis)

These results show significantly higher HDS rates for the isom-enhanced catalysts relative to KF756 alone.

These catalysts were subsequently tested on a feed spiked with a low level of organonitrogen to test the tolerance of the system to this poison.

**Example 25 (High HDS Activity)**

The distillate feed of Example 20 was spiked to 507 wppm S with 4.6-diethylidibenzothioophene and 11 wppm N with tetrahydroquinoline (THQ) and
processed over the catalyst system of Example 22. Results are presented in Table 3.

**Example 26 (High HDS Activity)**

The distillate feed of Example 25 was processed over the catalyst system of Example 23. Results are presented in Table 8.

**Example 27 (Comparative)**

A reactor was charged with 3.96 g (5.1 cc) of 14/35 mesh KF756 and was used to process the feed of Example 25. Results are presented in Table 8.

**Table 8**

**Comparison of HDS for KF756 and KF756 + Solid Acid**

<table>
<thead>
<tr>
<th>Example</th>
<th>Catalyst(s)</th>
<th>Product S (wppm)</th>
<th>RVA* For HDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>8:1 (w/w) KF756 + Pt/ECR-32(13)</td>
<td>185</td>
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<tr>
<td>8</td>
<td>KF756</td>
<td>295</td>
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325°C, 300 psig, 2.0 LHSV (over KF756) and 1000 SCF/B H₂

*Relative to KF756 alone (1.0 order HDS rate basis)
CLAIMS:

1. A two stage hydrodesulfurizing process for reducing the sulfur content of distillate feedstocks having a sulfur content greater than about 3,000 wppm, which process comprises:

   a) reacting said feedstream in a first hydrodesulfurization stage in the presence of a hydrogen-containing treat gas, said first hydrotreating stage containing one or more reaction zones, each reaction zone operated at hydrodesulfurizing conditions and in the presence of a hydrodesulfurization catalyst, thereby resulting in a liquid product stream having a sulfur content less than about 3,000 wppm;

   b) passing the liquid product stream of said first hydrodesulfurization stage to a separation zone where a vapor phase product stream and a liquid phase product stream are produced;

   c) reacting said liquid phase product stream of b) above in a second hydrodesulfurization stage in the presence of a hydrogen-containing treat gas said second hydrodesulfurization stage containing one or more reaction zones operated at hydrodesulfurization conditions wherein each reaction zone contains a bed of hydrotreating catalyst, thereby resulting in a liquid product stream having less than about 1,000 wppm sulfur;

   d) passing the liquid product stream of step c) above to a separation zone wherein a vapor phase stream and a liquid phase stream are produced;

   e) collecting both said vapor phase stream and said liquid phase stream; and

   wherein at least one of the reaction zones of at least one of said hydrodesulfurizing stages contains a bulk multimetallic catalyst comprised of at
least one Group VIII non-noble metal and at least two Group VIIB metals and wherein the ratio of Group VIIB metal to Group VIII non-noble metal is from about 10:1 to about 1:10.

2. The process of claim 1 wherein the Group VIII non-noble metal is selected from Ni and Co and the Group VIIB metals are selected from Mo and W.

3. The process of claim 1 wherein the bulk multimetallic is represented by the formula:

\[(X)_b (Mo)_c (W)_d O_x\]

wherein X is a Group VIII non-noble metal, and the molar ratio of b: (c+d) is 0.5/1 to 3/1.

4. The process of claim 3 wherein the molar ratio of b: (c+d) is 0.75/1 to 1.5/1.

5. The process of claim 3 wherein the molar ratio of c:d is preferably >0.01/1.

6. The process of claim 1 wherein the bulk multimetallic catalyst is essentially an amorphous material having a unique X-ray diffraction pattern showing crystalline peaks at d = 2.53 Angstroms and d = 1.70 Angstroms.

7. The process of claim 1 wherein the bulk multimetallic catalyst also contains an acid function.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER
   IPC(7) : C10G 45/04, 45/60, 29/04, 11/00, 11/02, 11/04; B01J 23/00
   US CL: Please See Extra Sheet.
   According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED
   Minimum documentation searched (classification system followed by classification symbols)
   Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched
   Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
   WEST
   search terms: hydrodesulfurization, hydrogenation, cobalt, nickel, molybdenum, tungsten

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
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<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tbody>
<tr>
<td>Y</td>
<td>US 3,925,197 A (VAN KLINKEN et al) 09 December 1975, col. 3, line 67 to col. 10, line 55.</td>
<td>1-7</td>
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<tr>
<td>A</td>
<td>US 3,861,005 A (STEINMETZ et al) 21 January 1975, col. 2, line 9 to col. 19, line 20.</td>
<td>1-7</td>
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<tr>
<td>A</td>
<td>US 3,850,746 A (ROBSON) 26 November 1974, col. 1, line 12 to col. 36, line 64.</td>
<td>1-7</td>
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<td>Y</td>
<td>US 4,014,925 A (FERLAZZO et al) 29 March 1977, col. 2, lines 10-24.</td>
<td>2-7</td>
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<tr>
<td>Y</td>
<td>US 4,808,563 A (VELENYI) 28 February 1989, col. 3, line 35 to col. 6, line 20.</td>
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</table>

☐ Further documents are listed in the continuation of Box C.  ☐ See patent family annex.

* Special categories of cited documents:
  "A" document defining the general state of the art which is not considered to be of particular relevance
  "R" earlier document published on or after the international filing date
  "A" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
  "P" document referring to an oral disclosure, use, exhibition or other means
  "F" document published prior to the international filing date but later than the priority date claimed
  "Y" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
  "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
  "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
  "A" document member of the same patent family

Date of the actual completion of the international search: 09 MARCH 2000
Date of mailing of the international search report: 11 APR 2000

Name and mailing address of the ISA/US Commissioner of Patents and Trademarks
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Washington, D.C. 20231
Facsimile No. (703) 305-3230
Authorized officer: MARK BELL
Telephone No. (703) 308-0661

Form PCT/ISA/210 (second sheet) (July 1992)*
**A. CLASSIFICATION OF SUBJECT MATTER:**

**US CL. :**

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<th>208/113, 121, 123, 124, 136, 137, 216R, 217, 295; 502/313, 315, 321</th>
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