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(54) REGENERATION OF SEVERELY DEACTIVATED REFORMING CATALYSTS

REGENERIERUNG VON SCHWER DESAKTIVIERTEN REFORMIERUNGSKATALYSATORS

REGENERATION DE CATALYSEURS DE REFORMAGE FORTEMENT DESACTIVES

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(56) References cited:
EP-A-0 142 352
EP-A-0 334 562

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This invention relates to regeneration of deactivated hydrocarbon processing catalysts comprising catalytic metals dispersed in a zeolite, particularly but not exclusively, reforming catalysts including one or more Group VIII metals in a large pore zeolite, such as platinum in zeolite L. The process of this invention is particularly effective for regenerating catalysts which have been more severely deactivated than is usual, for example, due to process upsets or to multi-cycle of on-oil regeneration operations. Such, catalysts regenerated in accordance with the present invention have been found to recover more of their activity and selectivity than if they were regenerated using conventional procedures. The general principles of the inventive process comprise (i) increasing the severity and time of the coke burn, (ii) using a source of high partial pressure chlorine in the oxychlorination step to redisperse catalytic metal e.g., platinum, (iii) improving chlorine removal by a wet, high temperature post treatment, at low pressure and (iv) enhancing reduction of catalytic metal by using high temperature at low pressure.

Catalytic reforming is a major petroleum refining process used to raise the octane rating of naphthas (C₈ to C₁₁ hydrocarbons) for gasoline blending. Catalytic reforming is also a principal source of aromatic chemicals, i.e., benzene, toluene and xylene, via conversion of paraffins and naphthenes to aromatics.

The principal reforming chemical reactions are dehydrogenation of cyclohexane to aromatics, dehydrocyclization of paraffins to aromatics, dehydroisomerization of alkylcyclopentanes to aromatics, isomerization of normal paraffins to branched paraffins, dealkylation of alkylbenzenes, and hydrocracking of paraffins to light hydrocarbons. The hydrocracking of paraffins to light hydrocarbons is undesirable and should be minimized because light hydrocarbons have low value.

Reforming is typically carried out at temperatures of 426° to 538°C (800°F to 1000°F), pressures of 0.34 to 20.7 MPa (50 to 3000 psi), hourly weight space velocities of 0.5 to 3.0, and in the presence of hydrogen at molar ratios to feed of 1 to 10. Other characteristics of reforming processes are well known in the art, and reference is made to US-A-4,914,068 for further teachings on this topic.

Catalysts currently used in commercial reformers include a Group VIII metal, such as platinum, or platinum plus a second catalytic metal, such as rhenium or iridium, dispersed on an alumina substrate. Typically, chlorine is incorporated on the alumina to add acid functionality. Alumina-based reforming catalysts are suitable for aromatizing C₈⁺ paraffins, but are less effective for aromatizing C₆ to C₈ paraffins because these catalysts hydrocrack more of the lighter paraffins to low value fuel gas than they convert to aromatics.

Conventional reforming catalysts are bifunctional, i.e., the catalysts enhance i) dehydrogenation and cyclization reactions on the catalytic metal sites; and ii) isomerization on separate strong acid sites in the catalyst. The undesirable hydrocracking reactions also occur on the acid sites.

Within the past few years reforming catalysts have been developed which have been discovered to be particularly effective for aromatizing the C₆ to C₈ paraffin components of naphtha. These catalysts are made using zeolite, rather than alumina, as the support for the catalytic metal. They are mono-functional and contain relatively few strong acid sites. Unlike conventional bifunctional catalysts, zeolite based catalysts accomplish dehydrogenation and cyclization reactions as well as isomerization on the dispersed metallic catalytic sites. Because these zeolite-based catalysts have few strong acid sites, undesirable hydrocracking reactions are repressed. Zeolites which are preferred for reforming catalysts are large pore zeolites i.e. zeolites with a 6 to 15 Angstrom pore diameter. Suitable large pore zeolites for purposes of the present invention include zeolite X, Y, and L, with zeolite L being the most preferred support for reforming catalysts, particularly wherein the catalytically active metal is platinum.

Compared to conventional alumina based reforming catalysts which are acidic and bifunctional, zeolite based catalysts such as zeolite L are non-acidic and monofunctional, which gives advantages in reforming light naphtha.

For production line-scale reforming processes, zeolite is preferably formed into aggregates, such as extrudates or tablets, typically 0.08 - 0.65 cm. in size; pressure drop across zeolite powder-packed reactor beds would be unacceptably high and power losses via elutriation in reactor off-gas would be excessive. For forming the zeolite into aggregates, oxides, such as alumina or silica, may be used as binder materials.

During reforming, catalysts deactivate due to accumulation of coke deposits in the catalyst and agglomeration of the fine catalytic metal particles dispersed in the catalyst into larger particles. Therefore, it is common practice to periodically regenerate reforming catalysts to recover their activity. Unexpectedly, however, it was found that it is more difficult to regenerate the previously described zeolite-based reforming catalysts than the conventional alumina-based catalysts. Platinum-on-zeolite L reforming catalysts are uniquely active and selective for aromatizing paraffinic C₆⁺, C₇⁺ and C₈⁺. Catalyst activity decreases progressively with time on oil due to build up of coke deposits on the catalyst and agglomeration of platinum, and although reactor inlet temperatures are periodically raised to compensate for
declining activity, periodically, the catalyst must be regenerated to recover its initial activity.

In general, such regeneration procedures include: 1) burning off the coke deposits using oxygen, 2) redispersing the active catalytic metals by oxyhalogenation using, for example, HCl, oxygen and water, 3) stripping excess halogen off the catalyst using first wet air, and then dry air as a stripping gas, and 4) reducing the dispersed catalytic metal using a reducing agent such as hydrogen.

For example, it is known that coke deposits may be removed from such catalysts by heating them in the presence of dilute oxygen at a flame-front temperature of 430°C to 540°C. This combustion may be preceded by flushing out residual hydrocarbon with inert gas, such as nitrogen. High temperature decoking however leads to loss of surface area of the supported metal particles and to removal of platinum from the zeolite channels, thereby resulting in loss of catalytic activity. Thus, after combustion, the catalyst is often subjected to oxychlorination by contact with oxygen and chlorine, or a chlorinated compound, such as CCl₄, at elevated temperatures.

In attempts to overcome problems experienced during regeneration of large-pore zeolite-based reforming catalysts, various procedures have been developed.

US-A-4,851,380 is directed to a process for regenerating sulfur contaminated reforming catalysts composed of a zeolite and Group VIII metal which involves intentionally agglomerating the Group VIII metal into large agglomerates and then treating the catalyst with a halogen acid gas to remove sulfur contaminants.

US-A-4,914,068 contains useful discussion on oxychlorination of decoked zeolite catalysts. However, the chlorine concentrations taught therein, at the specified total pressures, equate to extremely low chlorine or HCl partial pressures which do not give the advantages of the instant invention. Moreover, US-A-4,914,068 requires a reduction step practiced at a temperature which is lower than that of the chlorination or oxychlorination step, and discloses the benefits of cooling the catalyst in the presence of oxygen before the reduction.

EP-A-142352, corresponding to US-A-4,925,819, teaches improving the activity and selectivity of deactivated type L zeolite-based catalysts by dispersing the catalytic metal throughout the pores or channels of the zeolite. Such procedures adequately recover the activity of catalysts which are normally deactivated. However, it has been found that by use of such procedures, selectivity of regenerated catalysts are typically about 5% to about 10% lower than fresh catalysts. Even with diligent operation, reformers sometimes go into upset condition which severely deactivate the catalyst. Examples of upset conditions include loss of hydrogen flow or high temperature excursions due to excessive cracking reaction. Notwithstanding the disclosed effectiveness of the procedures of EP-A-142352 (US-A-4925819), these procedures do not fully recover the activity of severely deactivated catalysts. However, it has now been found that by incorporating certain critical modifications as listed above, the procedure of the present invention is capable of regenerating severely deactivated catalysts, and also, surprisingly, of achieving better selectivity with normally deactivated catalysts.

**SUMMARY OF THE INVENTION**

According to the present invention there is provided a process for regenerating a deactivated catalyst comprising a Group VIII catalytic metal and zeolite, which process comprises:

a) burning coke off the deactivated catalyst and converting the metal to agglomerated particles which are accessible to chlorine-containing gas in subsequent process step b) by contacting the catalyst for at least 6 hours with a gaseous stream comprising oxygen, inert gas and water under oxidation conditions comprising a temperature of from 400°C to 600°C to substantially decock the catalyst and agglomerate the catalyst to the extent that at least 80% by weight of the metal is agglomerated outside the channels of the zeolite as particles greater than 200 Å in size;

b) chlorinating and dispersing the metal of the substantially decoked catalyst by contacting the catalyst with a gaseous stream comprising water, a source of chlorine, oxygen and an inert gas under oxychlorination conditions comprising a temperature of from 450°C to 550°C and a partial pressure of chlorine derived from the source of chlorine, which is at least 207 Paa (0.03 psia), until the hydrogen chloride and/or chlorine which breaks through the catalyst reaches an HCl + Cl₂ partial pressure greater than 138 Paa (0.02 psia) to substantially completely chlorine-complex and disperse the metal;

c) removing at least some of the chlorine from the metal by contacting the chlorinated catalyst with a gaseous stream comprising water, oxygen, and an inert gas under chlorine removal conditions at a temperature of from 450°C to 550°C, and a total pressure below 690 kPa (100 psia), until the hydrogen chloride in the gaseous stream after contact with the catalyst falls to a partial pressure below 27.6 Paa (0.004 psia); and
d) reducing at least some of the chlorine-complexed metal in the catalyst to the metallic state by contacting the chlorinated catalyst from step (c) with a gaseous stream comprising inert gas and hydrogen under reducing conditions at a total pressure below 690 kPa (100 psia) and a temperature of from 350°C to 550°C to result in a regenerated catalyst comprising dispersed metal and zeolite.
The innovative modifications in the regeneration procedure of the present invention, over the previously identified regeneration procedures, include: (i) increased severity and time of coke burn, i.e., running the coke burn longer and at significantly more severe conditions; (ii) use of a source of chlorine capable of yielding chlorine at a partial pressure of at least 207 Pa (0.03 psia), such as elemental chlorine, i.e., Cl₂, HCl or other chlorine-containing agent to oxychlorinate and to redisperse the metal, preferably Pt; (iii) improved Cl⁻ removal after oxychlorination by operating at high temperature, and at low pressure, i.e., performing the postoxychlorination treatment at higher temperature to strip off more residual chlorine; and (iv) enhanced metal e.g., Pt reduction by operating at high reduction temperature to complete reduction of the metal e.g., platinum to the zero valent state and reducing at low pressure. In addition, the decoked catalyst may be reduced with hydrogen before step (b), in which case an inert purge step may be required after the hydrogen reduction and before oxychlorination in step (b). The improved regeneration procedure of the present invention may be conducted at a pressure of from sub-atmospheric to 2.07 MPa (300 psia), although steps (c) and (d) are conducted at low partial pressures to ensure appropriate stripping of chloride from the catalyst. The concentrations of the gases used depend upon the operating pressure and are adjusted accordingly to yield a desired range of partial pressure of the gas component wherein the partial pressure of gas A = the concentration of A x the total pressure.

Although not wishing to be bound by any particular theory, we believe that in practice of prior art regeneration techniques, the reactive gases in the oxychlorination step do not get access to all the catalytic metal e.g., Pt particles inside the zeolite channels in a severely deactivated catalyst. It is believed that increasing the severity of the coke burn step, as required by step (a) of the invention, permits the Group VIII catalytic metal to be accessible to subsequent treatment with chlorine-containing gas because of expulsion of the metal, e.g., of platinum, from within the microchannels of the zeolite. This step also clears the microchannels of foreign matter such as refractory (difficult to remove) coke which, if not removed, would interfere with reforming reactions. The metal, e.g., platinum, agglomerates outside channels of the zeolite into particles which have been found to be larger than with prior art procedures. The presence of a substantial proportion, e.g., greater than 80 or 90% by weight, of the metal in the form of particles greater than 200 Å in size has been found to indicate that the metal has been cleared from the microchannels and is accessible by the chlorine gas in step (b). Thus in accordance with the invention the zeolite has channels containing particles of Group VIII metal, and the temperature and the time of the oxidation conditions of step (a) are sufficient to remove a substantial amount of the particles of metal from inside to outside the channels in the zeolite, preferably to the extent that the metal outside the channels is in the form of agglomerated particles, having a size greater than about 200Å. The catalytic metal is preferably platinum, although references hereinafter to platinum should be taken as equally applicable to other Group VIII metals.

In step (b), chlorine at high concentration is required to decompose and redisperse these larger metal e.g., platinum, particles. With more aggressive chlorination it is necessary to enhance the post treat step (c) and platinum reduction step (d) to ensure adequate removal of residual chlorine and complete reduction of catalytic metal.

In addition to required steps (a), (b), (c) and (d), other process steps are optionally employed. Thus according to one embodiment, between steps (a) and (b) there is performed an intermediate reducing step (a') comprising contacting the substantially decoked catalyst with a gaseous stream comprising inert gas and hydrogen under reducing conditions for a time effective to reduce the metal component of the catalyst to the metallic state.

According to another embodiment, between steps (c) and (d) there is performed a purging step (c') comprising contacting the chlorinated catalyst from step (c) with an oxygen-free displacement gas flow for a time sufficient to purge oxygen therefrom prior to performing step (d).

**DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS**

The improved regeneration procedure of the present invention has four distinct steps: a) coke burn, b) oxychlorination, c) stripping, and d) hydrogen reduction.

(a) coke burn step

Step (a) is preferably performed at a temperature greater than 510°C and/or at an oxygen partial pressure not greater than 69 kPa (10 psia) and/or at a water partial pressure not greater than 10.3 kPa (1.5 psia) and/or at a total pressure of from sub-atmospheric to 2.07 MPa (300 psia). Preferably, the temperature is from 520°C - 600°C, more preferably 530°C - 550°C, and 540°C being most preferred. The coke burn step should ideally be conducted for a time longer than about 6 hours, e.g. 10 - 100 hours, more preferably 36 - 80 hours, with 48 - 75 hours being more preferred and 60 - 75 hours being most preferred. In this step, the partial pressure of water is preferably up to 5.2 kPa (0.75 psia), and is more preferably from 1.38 to 5.2 kPa (0.2 to 0.75 psia), and most preferably from 1.38 to 3.45 kPa (0.2 to 0.5 psia).

Preferably, step (a) is performed for a period of time until the gaseous stream after contact with the catalyst comprises a partial pressure of carbon dioxide of less than 69 Pa (0.01 psia), more preferably less than 27.6 Pa (0.004 psia). In a preferred embodiment, step (a) is performed in a first stage (a1) and a second stage (a2) wherein stage (a1)
is performed at a lower temperature e.g., 400 - 500°C, than stage (a2), e.g., 520-600°C.

(b) oxychlorination step

In this stage, the substantially deoked catalyst is treated with a second wet oxychlorination gas stream comprising chlorine from a source of chlorine yielding chlorine at a high partial pressure, preferably from 207 Pa to 20.7 kPa (0.03 to 3 psia), more preferably 345 Pa to 6.9 kPa (0.05 to 1.0 psia), and most preferably 690 Pa to 3.45 kPa (0.1 to 0.5 psia).

In a preferred embodiment, step (b) is performed at a temperature of from 480°C to 550°C and/or at an oxygen partial pressure not greater than 69 kPa (10 psia) and/or at a water partial pressure of not greater than 10.3 kPa (1.5 psia) and/or at a total pressure of from sub-atmospheric to 2.07 MPa (300 psia) and/or for at least two hours after HCl and/or Cl₂ breakthrough.

(c) wet post oxychlorination stripping step

In this stage, excess chlorine is removed from the chlorinated catalyst using a third wet gaseous stream, preferably at a temperature of from 480 to 520°C and/or at a water partial pressure not greater than 10.3 kPa (1.5 psia), more preferably from 345 Pa to 2.76 kPa (0.05 to 0.4 psia) and/or at an oxygen partial pressure of less than 31 kPa (4.5 psia), more preferably from 4.82 to 20.7 kPa (0.7 to 3 psia). The total pressure of step (c) is required to be low, i.e., below 690 kPa (100 psia) e.g., from sub-atmospheric to 448 kPa (65 psia), most preferably at about atmospheric pressure. The step is preferably performed for a time sufficient to effect a chloride concentration in offgas from the chlorinated catalyst of less than about 200 vppm at one atmosphere, which corresponds to a partial pressure of about 20.7 Pa (0.003 psia) more preferably at an offgas HCl concentration of less than 10.3 Pa (0.0015 psia).

(d) reduction step

In this stage, chlorinated catalyst from which excess chlorine has been removed is reduced by exposure to a fourth wet gaseous stream. Preferably step (d) is performed at a hydrogen partial pressure greater than 103 Pa (0.015 psia) e.g., from 103 Pa to 34.5 kPa (0.015 to 5 psia), more preferably from 1.03 to 34.5 kPa (0.15 to 5 psia) and/or a water partial pressure not greater than 10.3 kPa (1.5 psia), e.g., 345 Pa to 2.76 kPa (0.05 to 0.4 psia), and/or at a temperature of from 450°C to 520°C more preferably, 480° to 520°C. The total pressure of step (d) is required to be performed at a low total pressure, i.e., below 690 kPa (100 psia), more preferably from sub-atmospheric to 448 kPa (65 psia), most preferably at about atmospheric pressure. In a preferred embodiment, step (d) is performed under conditions such that the gaseous stream after contact with the catalyst comprises hydrogen chloride at a partial pressure of less than 51.7 Pa (0.0075 psia).

In addition to the foregoing, purge steps may be used between any of the previously described stages as desired or as deemed to be necessary from the standpoints of safety and technology. In this regard, an inert gas e.g., nitrogen purging step should ideally be used before reduction to remove oxygen from the reactor to avoid forming an explosive gas mixture. Also, the optional hydrogen reduction step (a') may be used after step (a) but before step (b) in which case the reduction with hydrogen should ideally be preceded and followed with a purging of an inert gas, such as nitrogen, before step b).

It will be understood that within each of steps (a) - (d) and the optional additional steps, there are several parameters (e.g., temperature, pressure, partial pressure, end points, concentrations) which may be varied within the required ranges in order to control the procedure and effect of the regeneration process. For ease of understanding of the scope of the invention, these parameters have been listed as "ranges of conditions" in Table 1. In this table the ranges of conditions given for each parameter of each process step are intended to be read independently of the ranges given for every other parameter. Those ranges indicated as suitable are intended to be understood as indicating those exemplary ranges which may be employed in performance of the invention, and not as any delineation of the scope of the invention. Further description of the preferred means of implementing the inventive process is provided hereinafter, and this should be read in conjunction with the "ranges of conditions" of Table 1.

a) Coke burn: The purpose of the coke burn step is to remove carbonaceous materials which accumulate on the catalyst during hydrocarbon processing off the catalyst by oxidizing it. The coke burn is best conducted in two phases - a primary coke burn (a1) followed by a secondary coke burn (a2). The primary coke burn is done at less severe conditions (lower temperature and lower inlet oxygen concentration) than the secondary coke burn. This is because thermal excursions to excessively high temperatures can occur if heavily coked deactivated catalyst were exposed to severe coke burn conditions directly.

As a practical matter, prior to regeneration of a deactivated catalyst, the reactors are purged to remove residual hydrocarbons left in the reactors when the hydroprocessing operation is terminated. This may be done by estab-
lishing a flow of hydrogen to the reactors and continuing hydrogen flow until hydrocarbon is displaced by hydrogen. Then flow of an inert gas such as nitrogen is established through the catalyst beds and maintained to purge the hydrogen. Linear gas velocity is maintained above about 15.2 cm/sec (0.5 FPS) for the purges and the coke burn and for all subsequent steps in the regeneration procedure. The inert gas flowing to the inlet of the reactors during the first stage of the coke burn step (a1) is heated to temperature using the feed preheater furnace. A gas stream comprising low concentrations of oxygen (Table 1) and an inert gas is used in the primary coke burn to avoid excessively high temperature, because of adiabatic coke burn condition. Under adiabatic coke burn, the increase in the flame front temperature depends solely on oxygen concentration regardless of the total pressure. High oxygen concentration can be used in primary coke burn when the coke burn is not carried out at the adiabatic condition, e.g. an external cooling to the reactor is available or a higher heat capacity inert gas other nitrogen is used. The primary coke burn step (a1) is preferably continued until the molar ratio of carbon dioxide to oxygen in the reactor outlet gas falls to about 3:1. The time to complete the primary coke burn phase varies with the concentration of coke on the catalyst, space velocity, and oxygen concentration in the reactor inlet gas and can take between five and fifty hours. The first gaseous stream in this stage of the process may be substantially dry or contain an amount of water at a partial pressure taught in Table 1.

The secondary coke burn step (a2) is initiated by raising inert gas temperature to above 510°C, and raising inert oxygen concentration to yield an oxygen partial pressure as taught in Table 1. Water is introduced to the reactor inlet gas stream to yield water partial pressure as taught in Table 1. In this regard, when the first stage burn is carried out at a lower water partial pressure than the second stage coke burn, it is preferred that the catalyst be saturated with the higher water partial pressure used in the second coke burn before increasing the temperature. Preferred temperatures, times, end points and pressures are as taught in Table 1. Lower and higher total pressures may be used when the concentrations of gases are adjusted accordingly to yield equivalent partial pressures of the gas component wherein the partial pressure of gas A = the concentration of A x the total pressure.

In addition to removing essentially all the coke from the catalyst, a second important purpose of the coke burn step is to expel the catalytic metals in zeolite microchannels to the outside surfaces of the catalyst and as a consequence the catalytic metal is agglomerated into particles which are typically larger than about 200 Å. The substantial removal of catalytic metal from inside the zeolite channels is assurred by conducting the secondary coke burn phase at a temperature above 510°C with water present for at least 6 hours. Although not wishing to be bound by any particular theory, it is believed that severe agglomeration conditions clears the zeolite microchannels of detritus such as refractory coke, and improves access to the catalytic metals for subsequent redispersion.

The effluent gas, i.e., the gas stream from the outlet of the vessel containing the catalyst, can be recycled during coke burn. If it is recycled then a fraction of the effluent gas must be purged and replaced with fresh inert gas to remove combustion products from the system. The purge rate is typically 5% to 30% of the recycle gas rate.

b) Oxychlorination: The purpose of the oxychlorination step is to redisperse the catalytic metals back into the zeolite microchannels by converting the metals to their chlorides. The metal chlorides are mobile and distribute facilely over the catalyst surfaces.

In the process of this invention, redispersion is accomplished by treating the decoked catalyst with a gas stream containing chlorine from a source of chlorine, oxygen, and water. Preferably the source of chlorine comprises organochloride compounds, elemental chlorine, and HCl. Preferred organochloride compounds comprise carbon tetrachloride, chloroform, methyl chloride, methylene chloride, dichloroethane, trichloroethane, and dichloroethylene, and mixtures thereof. Preferably the source of chlorine is Cl₂, HCl, or CCl₄, most preferably Cl₂. The oxychlorination step is conducted suitably at a total pressure at or above atmospheric pressure, as shown in Table 1. The concentrations of oxygen, water, and chlorine-containing compound injected at the reactor inlet are adjusted according to the reactor total pressure to yield the respective reactor inlet partial pressures, as shown in Table 1, wherein the partial pressure of gas A = the concentration of gas A x the total pressure.

The oxychlorination is continued at least until HCl and/or chlorine appears in the reactor outlet gas stream at a partial pressure greater than about 138 Paa (0.02 psia) which is referred to herein as the “breakthrough” of HCl and/or chlorine. After breakthrough, it is preferred to continue oxychlorination for two to four hours to make sure all the catalyst is thoroughly oxychlorinated.

c) Stripping: The purpose of stripping is to remove excess chlorine from the catalyst. Chlorine evolves during the subsequent hydrogen reduction step as HCl and induces agglomeration of the catalytic metal particles in proportion to HCl partial pressure. Accordingly, within limits, the more chlorine stripped from the catalyst prior to reduction, the higher will be the activity of the regenerated catalyst.

To improve the removal of chlorine, stripping is done at low pressure e.g., about atmospheric pressure. Total pressure and component partial pressures are as shown in Table 1. Stripping is continued until the HCl partial pressure in the reactor offgas falls below the levels shown in Table 1. This typically can take between about one and thirty hours. As previously mentioned, oxygen flow to the catalyst is terminated and the flow of wet inert gas is continued for a time to thoroughly purge oxygen from the system prior to introducing H₂.

To promote and enhance the efficiency of chlorine evolution, it is important that the stripping step be done at
as low a pressure as practicable. In most cases this is about atmospheric pressure; however, stripping can be enhanced by pulling vacuum on the reactors. Also, it is important to maintain a temperature of at least about 450°C, preferably within the ranges shown in Table 1. However, stripping may be carried out at higher reactor pressure with reduced stripping efficiency and therefore requires a longer time of stripping.

The effluent gas, i.e., the gas stream from the outlet of the vessel containing the catalyst, can be recycled. If it is recycled then a fraction of the effluent gas must be purged and replaced with fresh inert gas. The purge rate is typically 5% to 30% of the recycle gas rate. Ideally, though, this step is once through.

d) Reduction: The final step is reduction of catalytic metals, e.g., platinum, now dispersed uniformly across the surfaces of the catalyst as chlorides, to small zero valent metal clusters.

This is accomplished by exposing the stripped catalyst to hydrogen at a temperature greater than about 450°C, preferably within the range as shown in Table 1. Oxygen left in the stripping step is purged with inert gas, such as nitrogen, down below the explosion limit prior to introducing hydrogen. The reduction is accomplished with a gas stream containing hydrogen having a preferred partial pressure of hydrogen greater than 103 Paa (0.015 psia), preferably within the range shown in Table 1. The gas stream at the reactor inlet for the hydrogen reduction step also may contain water at a partial pressure as shown in Table 1, and the reduction is preferably conducted for a period of one to two hours. Although the exact reduction conditions used may vary, higher regenerated catalyst activity is achieved when the reduction temperature is over 450°C. Hydrogen feed rate is controlled such that the partial pressure HCl in reactor ofgas does not exceed 51.7 Paa (0.0075 psia). Preferably, the reduction step (d) is performed at a reactor total pressure of atmospheric to 448 Paa (65 psia), and preferably at about atmospheric. However, reactor pressure higher than 448 Paa (65 psia) may be used in this reduction step. For an easy operation, it is preferably to operate this reduction step at the same pressure as the previous stripping step.

In this hydrogen reduction step, hydrogen and inert gas may be recycled to the reactor inlet with a 10 to 30% purge. Preferably, this reduction step is carried out without recycle. In such embodiment, however, the catalyst has to be dried out after wet reduction in step (d) and before introducing hydrocarbon feed into the reactor with a drying gas comprising e.g., nitrogen or hydrogen, to a preferred water partial pressure of less than 27.6 Paa (0.004 psia), more preferably less than 20.7 Paa (0.003 psia), and most preferably less than 10.3 Paa (0.0015 psia). This drying step, therefore, is accomplished by reducing the partial pressure of water in the gas stream introduced at the inlet of the vessel containing the catalyst through the introduction of the drying gas so as to result in a reduced partial pressure of water in the gas at the outlet. Thus, the gas at the outlet is substantially devoid of water.

As previously mentioned, an optional hydrogen reduction step may be introduced after step (a) and before step (b).

As previously discussed, before hydrogen is introduced, however, oxygen remaining in the reactor after the coke burn must be purged from the reactor to preclude forming combustible mixtures of hydrogen and oxygen. In this stage, oxygen purging is preferably accomplished by cutting off oxygen flow and continuing nitrogen flow through the reactors.

For purposes of the present invention, feeding nitrogen for about 30 minutes to about two hours, with one hour being most preferred, at a temperature of about 510°C and a pressure within the range of between about atmospheric and 2.07 MPaa (300 psia) are suitable for reducing oxygen concentration to a safe level; and the nitrogen linear velocity is maintained in the catalyst beds at least about 15.2 cm/sec (0.5 FPS) to achieve adequate gas distribution and gas/solid contacting.

Hydrogen reduction step (a') is initiated by starting flow of hydrogen into the nitrogen flowing into the reactors. Conditions of hydrogen reduction can be varied over a wide range, as shown in Table 1.

After completing hydrogen reduction, hydrogen flow is terminated and inert gas flow is continued to purge hydrogen so that oxygen required in the next step can be safely introduced. The purge can be done over a wide range of conditions; for example, passing inert gas through the catalyst at about atmospheric to about 2.07 MPaa (300 psia) and about 450°C to about 550°C, preferably between about 450°C to about 530°C, above about 15.2 cm/sec (0.5 FPS) for about 30 minutes adequately clears hydrogen.
TABLE 1
RANGES OF CONDITIONS

<table>
<thead>
<tr>
<th>Components</th>
<th>Suitable</th>
<th>Preferable</th>
<th>More Preferable</th>
<th>Most Preferable</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary Coke Burn</strong>&lt;br&gt;(a1)&lt;br&gt;Oxygen</td>
<td>500-6000 ppm</td>
<td>1000-4000 ppm</td>
<td>0-0.1 psia</td>
<td>50-200 psia</td>
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<td>Water</td>
<td>0-1.5 psia</td>
<td>0-0.75 psia</td>
<td>0-0.75 psia</td>
<td>14.7-200 psia</td>
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<td>Pressure</td>
<td>14.7-300 psia</td>
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<td>400-450°C</td>
<td>14.7 psia</td>
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<td>Temperature</td>
<td>400-500°C</td>
<td>400-480°C</td>
<td>400-450°C</td>
<td>400-500°C</td>
</tr>
<tr>
<td><strong>Secondary Coke Burn</strong>&lt;br&gt;(a2)&lt;br&gt;Oxygen</td>
<td>up to 10 psia</td>
<td>0.1-5 psia</td>
<td>0.2-5 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Water</td>
<td>up to 1.5 psia</td>
<td>up to 0.75 psia</td>
<td>0.2-5 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Pressure</td>
<td>14.7-300 psia</td>
<td>14.7-200 psia</td>
<td>0.2-5 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Time</td>
<td>&gt;6 hours</td>
<td>36-80 hours</td>
<td>48-75 hours</td>
<td>60-75 hours</td>
</tr>
<tr>
<td>End Point</td>
<td>CO2&lt;0.01 psia</td>
<td>CO2&lt;0.004 psia</td>
<td>HCl and Cl2</td>
<td>HCl and Cl2</td>
</tr>
<tr>
<td>Temperature</td>
<td>&gt;510°C</td>
<td>520-600°C</td>
<td>530-550°C</td>
<td>530-550°C</td>
</tr>
<tr>
<td><strong>Oxychlorination</strong>&lt;br&gt;(b)&lt;br&gt;Oxygen</td>
<td>up to 10 psia</td>
<td>1-5 psia</td>
<td>2-4 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Water</td>
<td>up to 1.5 psia</td>
<td>up to 0.75 psia</td>
<td>0.2-0.5 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Chlorine</td>
<td>&gt;0.03 psia</td>
<td>0.03-3 psia</td>
<td>0.1-0.5 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Source of Cl</td>
<td>Chlorohydrocarbons</td>
<td>HCl, Cl2, and</td>
<td>Cl2</td>
<td>HCl and Cl2</td>
</tr>
<tr>
<td>Pressure</td>
<td>&gt;14.7 psia</td>
<td>14.7 psia</td>
<td>14.7 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Temperature</td>
<td>480-550°C</td>
<td>500-530°C</td>
<td>530-550°C</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Breakthrough</td>
<td>HCl or Cl2&gt;0.02 psia</td>
<td>HCl&lt;0.004 psia</td>
<td>HCl&lt;0.0015 psia</td>
<td>510°C</td>
</tr>
<tr>
<td><strong>Stripping</strong>&lt;br&gt;(c)&lt;br&gt;Oxygen</td>
<td>&lt;4.5 psia</td>
<td>&lt;3 psia</td>
<td>0.7-3 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Water</td>
<td>up to 1.5 psia</td>
<td>0-0.75 psia</td>
<td>0.05-0.4 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Pressure</td>
<td>14.7-65 psia</td>
<td>14.7 psia</td>
<td>0-0.4 psia</td>
<td>530-550°C</td>
</tr>
<tr>
<td>Temperature</td>
<td>&gt;450°C</td>
<td>450-530°C</td>
<td>480-520°C</td>
<td>510°C</td>
</tr>
<tr>
<td>End Point</td>
<td>HCl&lt;0.004 psia</td>
<td>HCl&lt;0.003 psia</td>
<td>HCl&lt;0.0015 psia</td>
<td>510°C</td>
</tr>
<tr>
<td><strong>Reduction</strong>&lt;br&gt;(d)&lt;br&gt;Hydrogen</td>
<td>&gt;0.015 psia</td>
<td>0.015-14.7 psia</td>
<td>0.15-5 psia</td>
<td>510°C</td>
</tr>
<tr>
<td>Water</td>
<td>0-1.5 psia</td>
<td>up to 0.75 psia</td>
<td>0.4-1 psia</td>
<td>510°C</td>
</tr>
<tr>
<td>Pressure</td>
<td>14.7-65 psia</td>
<td>14.7 psia</td>
<td>0.05-0.4 psia</td>
<td>510°C</td>
</tr>
<tr>
<td>Temperature</td>
<td>&gt;450 °C</td>
<td>450-530°C</td>
<td>480-520°C</td>
<td>510°C</td>
</tr>
<tr>
<td>HCl Release</td>
<td>&lt;0.0075 psia</td>
<td>510°C</td>
<td>480-520°C</td>
<td>510°C</td>
</tr>
<tr>
<td><strong>Option Reduction After coke burn</strong>&lt;br&gt;(a')&lt;br&gt;Hydrogen</td>
<td>&gt;0.015 psia</td>
<td>0.015-14.7 psia</td>
<td>0.15-5 psia</td>
<td>510°C</td>
</tr>
<tr>
<td>Water</td>
<td>0-1.5 psia</td>
<td>0-0.75 psia</td>
<td>0.2-0.5 psia</td>
<td>510°C</td>
</tr>
<tr>
<td>Pressure</td>
<td>14.7-300 psia</td>
<td>14.7-200 psia</td>
<td>14.7 psia</td>
<td>510°C</td>
</tr>
<tr>
<td>Temperature</td>
<td>300-500°C</td>
<td>450-530°C</td>
<td>480-520°C</td>
<td>510°C</td>
</tr>
</tbody>
</table>
The effluent gas, i.e., the gas stream from the outlet of the vessel containing the catalyst, can be recycled. If it is recycled then a fraction of the effluent gas must be purged and replaced with fresh inert gas. The purge rate is typically 5% to 30% of the recycle gas rate.

For purposes of the present invention, preferred catalysts are monofunctional catalysts based on large pore zeolite crystals bound together by a binder which is preferably kaolin, silica, or most preferably alumina. The large pore zeolite is preferably zeolite L and more preferably includes exchangeable cations at least a portion of which are selected from lithium, sodium, potassium, rubidium, cesium, magnesium, calcium, strontium, barium, and mixtures of any two or more thereof. More preferably the exchangeable cation is potassium or barium, and most preferably potassium.

The Group VIII catalytic metals are preferably noble metals e.g., platinum, palladium, rhodium, iridium, ruthenium and osmium, and most preferably platinum. The amount of the Group VIII catalytic metal present is preferably from 0.05% to 6% by weight.

The catalytic metal may further comprise a Group VII B metal, such as rhenium. The Groups identified herein are as presented in the Periodic Table in the Handbook of Chemistry & Physics, published by the Chemical Rubber Company, 48th edition.

In a preferred embodiment, the base of the catalyst is approximately 70% zeolite L and 30% alumina by weight and is formed into 0.16 cm extrudates. For purposes of the present invention, however, binders other than alumina, such as silica, kaolin and clay may be used; and formed shapes, other than extrudates, such as spheres, tablets, pellets and extrudates may also be used. The formed catalyst is preferably loaded with platinum by ion exchange so that the metal loading is preferably between 0.5% and 0.9% by weight. Representative publications disclosing details of catalyst preparation, characterization, and manufacture include US-A-3,216,789 which describes zeolite L; US-A-4,104,320; 4,416,806; and, 4,568,656 which describe processes of ion exchange of platinum into zeolite L; and US-A-4,595,668; 4,595,669; and 4,595,670 which describe preferred reforming catalysts comprising platinum on zeolite L which has finely dispersed platinum.

The activity of a catalyst is a measure of its ability to convert feed to products. While a catalyst may have a high activity, the products formed may not be necessarily the desired products. The term “selectivity” is a measure of the ability of the catalyst to convert feed to desired products. Activity maintenance concerns the ability of the catalyst to maintain activity over time at conversion conditions.

The decline in catalyst activity at conversion conditions is believed to be due primarily to crystal growth or agglomeration of the noble metal particles to about the size of the zeolite channels (about 12 Å) and secondarily to the formation of coke on the external and internal surfaces of the catalyst. At the same noble metal loading, catalysts containing particles or crystals of noble metals of a size greater than that disclosed above are less active and less selective than catalysts containing the smaller particles. Moreover, agglomerated metal particles block zeolite channels. Coke formation, probably due to complex condensation and dehydrogenation reactions, results in the shedding of the noble metal from the reaction mixture, thereby limiting the catalytic sites available for promoting reactions.

As catalytic activity declines by virtue of agglomeration and coke formation, yield of desirable products will decline, and depending upon the economics of the operation a process employing the catalyst will have to be interrupted and catalyst activity restored to its initial value. Generally, catalytic activity can be maintained by increasing temperature, but there are limits beyond which the temperature cannot be increased, for example, temperatures which will change the nature of the zeolite or lead to undesirable side reactions or exceed reactor material limitations. Particularly for zeolite based catalysts, temperatures higher than 538°C or 594°C (1000°F or 1100°F) are preferably avoided.

Catalyst activity will decline faster over time as severity increases. Factors which increase severity include: increasing temperature, reducing hydrogen to oil mole ratio, reducing hydrogen partial pressure, reducing total pressure and temperature, increasing feed rate per volume of catalyst (space velocity), and increasing type of hydrocarbon in the feed.

In the measurement of activity maintenance all variables are fixed and only the catalyst differs. Thus, an activity of one catalyst over a period of time can be directly compared to the activity of another catalyst over the same time period where feed, hydrogen to oil ratios, pressures, etc., are constant.

One activity test utilized for purposes of the present application is to pass a feedstream comprising about 40 wt. % normal hexane and 60% isohexanes over the catalyst at the following conditions: 510°C, 1.0 weight hourly space velocity (WHSV), 839 kPa (121.7 psia), 4.25 H₂/oil molar ratio. Benzene yield as a percent of feed after 46 hours is a good measure of catalyst activity.

Although the regeneration process herein may apply to deactivated catalysts at any stage of deactivation, it is particularly suitable for those which are severely deactivated.

EXAMPLES

The following examples illustrate the invention. In all examples, parts and percentages are given by weight for solids and liquids and by volume for gas compositions, and temperatures in degrees Centigrade unless otherwise noted. Examples 1 through 18 were performed at one atmospheric total pressure. In all Examples, the HCl level in the effluent
gas after the stripping step (c) and after the reduction step (d) was less than 27.6 Paa (0.004 psia) and in most cases less than 2.76 Paa (0.0004 psia) partial pressure, as measured or calculated.

Example 1

A 0.64 wt % Pt on KL-zeolite extruded catalyst was used to aromatize a light naphtha feed. The flow of hydrogen to the reactor was cut off and the catalyst was exposed to pure hydrocarbon feed for about 5 hours at reforming conditions. The catalyst was thus deactivated, i.e., coked up severely, as indicated by accumulation of about 15 wt. % by weight of coke in contrast to a normally deactivated catalyst which typically contains about 1 wt. % coke.

Two grams of the severely deactivated catalyst were regenerated by the following standard HCl + O2 redispersion procedure:

First coke deposit was removed by burning (a1) at 450°C with 20 vol. % O2 in helium at 500 cc/min. for 30 minutes; then temperature was increased to 510°C and water at 2.2 vol. % was added, and the catalyst was subjected to these conditions (a2) for 2 hours. Next HCl was added to the gas stream to give 0.16 vol. % and at the same time oxygen concentration was reduced to 10 vol. %, resulting in a low approaching zero Cl2 partial pressure. The total gas flow rate was maintained at 500 cc/min. The oxchlorination treatment (b) was performed for 2.5 hours with a breakthrough HCl + Cl2 partial pressure of 166 Paa (0.024 psia). After the previously described oxchlorination procedure, the catalyst was treated (c) with 2.2 vol. % water, 10 vol. % O2 and 87.8 vol. % helium for 1 hour. Oxygen was purged out in about 10 minutes with wet helium containing about 2% water vol. %. Hydrogen reduction (d) was conducted at 510°C with 10 vol. % H2, and 2.2 vol. % H2O and 87.8 vol. % helium at a flow rate of 500 cc/min. for 1.5 hours. The regenerated catalyst was designated as Catalyst A.

Example 2

The coked catalyst used in Example 1 was regenerated with the regeneration procedure as described in Example 1 except that during the second stage (a2) of the coke burn procedure, the temperature was increased to about 530°C which was thereafter employed and the treat time was extended to about 6 hours; also during oxchlorination, 0.82 vol% Cl2 instead of HCl was employed, [Cl2 partial pressure and breakthrough HCl and Cl2 partial pressure each approx. 834 Paa (0.121 psia)]. The regenerated catalyst regenerated in accordance with the present invention was designated as Catalyst B.

Example 3

The above two catalysts were evaluated using 3-methylpentane feed. The reaction was carried out at 510°C at a space velocity of 20 w/w/hr, H2/oil-6 (6 molar ratio of H2 to oil), and 724 Paa (105 psia). Table 2 shows the catalyst performance of Catalyst A and B. Catalyst B which was regenerated by a process in accordance with the present invention shows a higher benzene yield and selectivity than Catalyst A.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wt% Benzene @ 10 Hr.</th>
<th>Wt% Benzene @ 25 Hr.</th>
<th>Selectivity @ 25 Hr.</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>24.5</td>
<td>20.2</td>
<td>67.3</td>
</tr>
<tr>
<td>B</td>
<td>26.3</td>
<td>22.6</td>
<td>68.8</td>
</tr>
</tbody>
</table>

Example 4

A 0.85 wt % Pt on KL-zeolite catalyst was deactivated during aromatization of a light naphtha feed. One gram of this catalyst was regenerated with the procedure of Example 1, except that after the second coke burn, the catalyst was reduced a’ at 510°C with 20 vol. % H2 and 2.2 vol. % water for 1 hour before an HCl + O2 oxchlorination. The HCl concentration was 0.32 vol. %, (breakthrough partial pressure approx. 324 Paa (0.047 psia); Cl2 partial pressure approx. zero). The regenerated catalyst was designated as Catalyst C.

Example 5

The coked catalyst used in Example 4 was regenerated with the same procedure as Example 2 except that in the
second coke burn, the burn time was increased to 22 hours and there was a wet hydrogen reduction (a') before the \( \text{Cl}_2 + \text{O}_2 \) oxychlorination, (\( \text{Cl}_2 \) and breakthrough partial pressure (pp) as reported in Example 2). The regenerated catalyst was designated as Catalyst D.

**Example 6**

The above two catalysts were evaluated using 3-methylpentane feed. The reaction was carried out at 510°C at a space velocity of 20 w/w/hr, \( \text{H}_2/\text{oil-6} \), and 724 Paa (105 psia). Table 3 shows the catalyst performance of Catalyst C and D. Catalyst D which was regenerated by the instant invention shows better catalyst performance than Catalyst C.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wt% Benzene @ 10 Hr</th>
<th>Wt% Benzene @ 25 Hr</th>
<th>Selectivity @ 25 Hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>22.2</td>
<td>18.3</td>
<td>69.6</td>
</tr>
<tr>
<td>D</td>
<td>25.2</td>
<td>20.8</td>
<td>69.8</td>
</tr>
</tbody>
</table>

**Example 7**

A catalyst was deactivated through 6 cycles of on-oil and regeneration operations in the Catalyst Treating Unit. One gram of this catalyst was regenerated using exactly the same procedure as Example 1 (\( \text{Cl}_2 \) and breakthrough pp as in Example 1). The regenerated catalyst was designated as Catalyst E.

**Example 8**

The deactivated catalyst used in Example 7 was employed in this example. Four grams of this catalyst was regenerated using exactly the same procedure as Example 2 except that the time of the coke burn step was extended to 69 hours (\( \text{Cl}_2 \) and breakthrough pp as in Example 2). The regenerated catalyst was designated as Catalyst F.

**Example 9**

The above two catalysts were evaluated using 3-methylpentane feed. The reaction was carried out at 510°C at a space velocity of 20 w/w/hr, \( \text{H}_2/\text{oil-6} \), and 724 Paa (105 psia). Table 4 shows the catalyst performance of Catalyst E and F. Catalyst F which was regenerated by the instant invention shows a better catalyst performance than Catalyst E.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wt% Benzene @ 10 Hr</th>
<th>Wt% Benzene @ 25 Hr</th>
<th>Selectivity @ 25 Hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>E</td>
<td>19.6</td>
<td>16.0</td>
<td>67.6</td>
</tr>
<tr>
<td>F</td>
<td>21.5</td>
<td>18.8</td>
<td>64.6</td>
</tr>
</tbody>
</table>

Advantages of the present invention in the regeneration of Pt on KL-zeolite include the externalization of the Pt particles to the outside surface of the zeolite crystallites during high temperature coke burn which appears to open up the zeolite channels for Pt redistribution in the oxychlorination step. The redispersion of large Pt particles which are larger than about 200 Å requires high concentration of chlorine to facilitate the formation of mobile Pt chloride species. Although not wishing to be bound by any particular theory, it is believed that this redistribution of external Pt into zeolite channels may promote highly dispersed Pt clusters which reside near the channel mouths thereby yielding better catalyst performance.

The following examples using fresh catalysts give some support to the above interpretation.

**Example 10**

A sample of about 3 grams of a 0.85 wt % Pt/KL-zeolite catalyst was subjected to a 510°C calcination of 20 vol. %
O₂, 2.2 vol. % H₂O and 77.8 vol. % helium at a flow rate of 500 cc/min. for 2 hours. Less than 10% of the Pt formed particles had a size greater than about 200 Å. Substantial amount of Pt remained inside the zeolite channels as large and small particles. This catalyst was designated at Catalyst G.

**Example 11**

Ten grams of the fresh catalyst used in Example 10 were calcined at 530°C in 20 vol. % O₂, 2.2 vol. % H₂O, and 77.8 vol. % helium at a flow rate of 500 cc/min. for 64 hours. Substantially all the Pt was removed from the zeolite channels to the external surface of the zeolite crystallites as particles larger than about 200 Å. This catalyst was designated as Catalyst H.

**Example 12**

Two grams each of Catalysts G and H were regenerated using the same regeneration procedure as that in Example 1 except that in oxychlorination higher HCl concentration (0.58 vol. %) was used, [breakthrough pp approx. 586 Paa (0.085 psia); Cl₂ pp approx. zero]. The regenerated catalysts were designated as Catalysts I and J, respectively.

**Example 13**

Two grams each of Catalyst G and H were regenerated using the same procedure as in Example 12 except that HCl concentration in oxychlorination was increased to 1.08 vol. %, [breakthrough pp approx. 1.1 kPaa (0.159 psia); Cl₂ pp approx. zero]. The regenerated catalysts were designated as Catalysts K and L respectively. Catalysts I, J, K, and L were evaluated by 3-methylpentane aromatization and hydrogen chemisorption as described in Example 3. The results are in Table 5. Higher concentrations of HCl produces higher Cl₂ concentration via the Deacon reaction:

\[
4 \text{HCl} + \text{O}_2 = 2 \text{Cl}_2 + 2 \text{H}_2\text{O}.
\]

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>H/Pt</th>
<th>Wt% Benzene @ 25 Hr</th>
<th>Bz Selectivity @ 25 Hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh</td>
<td>0.78</td>
<td>19.4</td>
<td>68.2</td>
</tr>
<tr>
<td>I</td>
<td>0.59</td>
<td>22.2</td>
<td>67.6</td>
</tr>
<tr>
<td>J</td>
<td>0.37</td>
<td>18.6</td>
<td>68.7</td>
</tr>
<tr>
<td>K</td>
<td>0.71</td>
<td>22.9</td>
<td>69.3</td>
</tr>
<tr>
<td>L</td>
<td>0.59</td>
<td>21.2</td>
<td>68.5</td>
</tr>
</tbody>
</table>

Table 5 indicates that higher HCl concentration in the HCl + O₂ oxychlorination improves platinum dispersion for the 510°C calcined catalysts (Catalysts I and K) from 0.59 to 0.71 and 530°C calcined catalysts (Catalysts J and L) from 0.37 to 0.59. Catalyst performance is also affected, especially in the case of catalysts J and L.

**Example 14**

Two grams each of Catalysts G and H were regenerated using the same regeneration procedure as that in Example 2 in which oxychlorination was carried out with 0.82 vol. % Cl₂ [breakthrough and Cl₂ pp as in Example 2]. The regenerated catalysts were designated as Catalysts M and N respectively. Catalysts M and N were evaluated by 3-methylpentane aromatization and hydrogen chemisorption [as in Example 3]. The results are in Table 6.
Table 6 indicates that high Cl₂ concentration completely redisperses the particles of Pt which are 200 Å formed during the 530°C calcination. Additionally, the 530°C calcined catalyst (N) showed a much higher activity than the fresh and the 510°C calcined catalyst (M) after Cl₂ + O₂ oxychlorination. Although not wishing to be bound by any particular theory, this suggests that redispersion of the large Pt particles on the external surface of the zeolite crystallites (530°C calcined catalyst (N)) leads to highly dispersed Pt clusters in the zeolite channels but close to the channel mouths compared to the case of redispersing agglomerated Pt particles inside zeolite channels (510°C calcined catalyst (M)).

Example 15

A 0.84 wt % Pt on KL-zeolite extruded catalyst was used to aromatize a light naphtha feed. The catalyst was severely deactivated and heavily coked. The color of the catalyst was black. Three grams of the catalyst were regenerated by the following HCl + O₂ redispersion procedure. First coke deposit was removed by burning at 450°C with 10 vol. % O₂ in helium at 1000 cc/min. for 1 hour. Then temperature was increased to 510°C and held at 510°C for 12 hours. Next water and HCl were added to the gas stream to give 3.3 vol. % water and 0.33 vol. % HCl, and at the same time oxygen concentration was still maintained at 10 vol. %. The total gas flow rate was maintained at 1000 cc/min. The oxychlorination treatment (b) was for 6 hours; [breakthrough pp 338 Paa (0.049 psia); Cl₂ pp approx. zero]. The catalyst was then cooled to 345°C under substantially same gas environment and was treated with 3.3 vol. % water, 10 vol. % O₂ and 87.8 vol. % nitrogen for 1 hour. Oxygen was purged out in about 10 minutes with nitrogen. Hydrogen reduction was conducted at 345°C with 10 vol. % H₂ and 90 vol. % helium at a flow rate of 1000 cc/min for 1 hour. The regenerated catalyst was designated as Catalyst O.

Example 16

The coked catalyst used in Example 15 was regenerated with the regeneration procedure in accordance with the present invention. The coke was first removed by burning (a1) at 445°C with 10 vol. % oxygen in nitrogen for 1 hour. The temperature was raised to 530°C under 20% O₂, 2.2% H₂O and balanced helium. The catalyst temperature was held (a2) at 530°C for 69 hours before it was cooled to 510°C. During oxychlorination (b), catalyst was treated with a gas mixture of 2.2%H₂O, 20%O₂, 1.5% Cl₂, and balanced He for 2.5 hours at 510°C, [Cl₂ and breakthrough pp = 1.52 kPaa (0.221 psia)]. After the oxychlorination, the catalyst was treated (c) with 2.2 vol. % water, 10 vol. % O₂, and 87.8 vol. % helium at 510°C for 1 hour. Oxygen was purged in about 10 minutes with wet helium. Hydrogen reduction (d) was conducted at 510°C with 20% H₂, 2.2 vol.% water and 87.8 vol. % helium at a flow rate of 500 cc/min. for 1.5 hours. The regenerated catalyst was designated Catalyst P.

Example 17

Catalysts O and P and fresh catalyst were evaluated by an aromatization reaction with a mixed feed of 60 wt% 3-methyl pentane and 40 wt% n-hexane. The reaction was carried out at 510°C at a space velocity of 8 w/w/hr. H₂/oil-2.5, and 724 Paa (107 psia). Table 7 shows the catalyst performance of fresh catalyst and Catalysts O and P. Catalyst P which was regenerated by the current invention shows a better catalyst performance than Catalyst O and fresh catalyst.

Table 7

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wt% Benzene @ 46 Hr</th>
<th>Bz Selectivity @ 46 Hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh</td>
<td>37</td>
<td>76</td>
</tr>
<tr>
<td>O</td>
<td>33</td>
<td>65</td>
</tr>
<tr>
<td>P</td>
<td>40</td>
<td>75</td>
</tr>
</tbody>
</table>
Example 18

A 0.85 wt% Pt on KL-zeolite catalyst was used to aromatize a light naphtha feed. The catalyst was normally deactivated in that it contained about 1-2% coke. The catalyst was regenerated by the conventional procedure as described in Example 15. This catalyst is designated as Catalyst Q.

In parallel, three grams of the catalyst were regenerated by the regeneration procedure in accordance with the present invention as described in Example 16. This catalyst is designated as Catalyst R. Catalysts Q and R were evaluated under the aromatization test condition described in Example 17. Table 8 indicates that the regeneration procedure in accordance with the present invention achieves higher activity and selectivity than a conventional procedure used to regenerate a normally deactivated catalyst.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wt% Benzene @ 46 Hr</th>
<th>Bz Selectivity @ 46 Hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>Q</td>
<td>37</td>
<td>72</td>
</tr>
<tr>
<td>R</td>
<td>40</td>
<td>75</td>
</tr>
</tbody>
</table>

Example 19

A 0.85 % wt. % Pt on formed KL-zeolite catalyst was used to aromatize a light naphtha feed. The catalyst was normally deactivated, i.e., the catalyst contained about 1-2% coke. The catalyst was regenerated using the following procedure:

The coke was removed by coke burn (a1) at 450°C for 10 hours with 0.2 vol. % oxygen in nitrogen under 1.28 MPaa (185 psia) total pressure and 15.2 cm/sec (0.5 FPS) linear velocity. The coke burn (a2) was further carried out at 510°C for 16 hours with 3.5 vol. % oxygen. After oxygen was purged out, the catalyst was reduced (a') at 510°C with 50% hydrogen for 1 hour. After hydrogen was purged out, the catalyst was then pre-conditioned for oxychlorination with 0.89 vol. % oxygen and 0.87 vol. % water vapor at 510°C under 1.17 MPaa (170 psia) total pressure. HCl was added to the stream to a concentration of 0.15 vol. % for 4.5 hours. After oxygen was purged out, the catalyst was then treated with 5 vol. % oxygen and 0.84 vol. % water vapor at 345°C under atmospheric pressure for 8 hours. Water vapor was removed from the system, and the catalyst was further treated at 345°C for 2 hours. After oxygen was purged out from the system, the catalyst was reduced at 345°C with 7 vol. % hydrogen for 6 hours. The catalyst is designated as Catalyst S.

A batch of the similar normally deactivated 0.85% Pt/KL-zeolite catalyst was regenerated by a regeneration procedure in accordance with the present invention. The coke was removed by coke burn (a1) at 450°C for 10 hours with 0.2 vol% oxygen in nitrogen under 1.17 MPaa (185 psia) total pressure and 15.2 cm/sec (0.5 FPS) velocity. The coke burn was further carried out (a2) at 530°C for 69 hours with 1.6 vol. % oxygen and 0.2 vol. % water vapor. The catalyst was then treated (b) at 510°C with 0.12 vol. % oxygen, 0.12 vol. % Cl₂, and 0.2 vol. % H₂O under 1.17 MPaa (185 psia) total pressure. After oxygen was purged out, the catalyst was then treated with 10.5 vol. % oxygen and 2.2 vol. % water vapor at 510°C under atmospheric pressure for 10 hours; to an HCl partial pressure of 45.5 Paa (0.0066 psia). After oxygen was purged out from the system, the catalyst was reduced (d) at 510°C with 5.8 vol. % hydrogen and 2.2 vol. % water vapor for 1.5 hours at atmospheric pressure, to an HCl pp of 15.2 Paa (0.0022 psia). The catalyst is designated as Catalyst T.

The activities of Catalysts S and T in light naphtha aromatization under the following conditions:

<table>
<thead>
<tr>
<th>Feedstream</th>
<th>Conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed C₅s</td>
<td>1.8 WHSV</td>
</tr>
<tr>
<td>60% nC₆</td>
<td>2 H₂/Oil Molar Ratio</td>
</tr>
<tr>
<td>40% iC₆</td>
<td>105 psia</td>
</tr>
<tr>
<td>505°C</td>
<td>505°C</td>
</tr>
</tbody>
</table>
The normally deactivated catalyst regenerated in accordance with the present invention has higher activity and selectivity than the catalyst regenerated by the HCl procedure.

The catalysts regenerated by the process of this invention may be employed as reforming catalysts using reforming conditions which are well known in the art.

In summary, the present invention is seen to provide a method which is particularly effective for regenerating severely deactivated reforming catalysts, preferably extrudates of potassium-exchanged zeolite L containing platinum, so as to substantially restore catalyst activity and activity maintenance to the catalyst. The regeneration procedure of the present invention, however, is also useful for regenerating normal deactivated reforming catalysts.

Claims

1. A process for regenerating a deactivated catalyst comprising a Group VIII catalytic metal and zeolite, which process comprises:

a) burning coke off the deactivated catalyst and converting the metal to agglomerated particles which are accessible to chlorine-containing gas in subsequent process step b) by contacting the catalyst for at least 6 hours with a gaseous stream comprising oxygen, inert gas and water under oxidation conditions comprising a temperature of from 400°C to 600°C to substantially decoke the catalyst and agglomerate the catalyst to the extent that at least 80% by weight of the metal is agglomerated outside the channels of the zeolite as particles greater than 200 Angstrom in size;

b) chlorinating and dispersing the metal of the substantially decoked catalyst by contacting the catalyst with a gaseous stream comprising water, a source of chlorine, oxygen and an inert gas under oxychlorination conditions comprising a temperature of from 450°C to 550°C and a partial pressure of chlorine derived from the source of chlorine which is at least 207 Paa (0.03 psia), until the hydrogen chloride and/or chlorine which breaks through the catalyst reaches an HCl + Cl2 partial pressure greater than 138 Paa (0.02 psia) to substantially completely chlorine-complex and disperse the metal;

c) removing at least some of the chlorine from the metal by contacting the chlorinated catalyst with a gaseous stream comprising water, oxygen, and an inert gas under chlorine removal conditions at a temperature of from 450°C to 550°C, and a total pressure below 690 kPaa (100 psia), until the hydrogen chloride in the gaseous stream after contact with the catalyst falls to a partial pressure below 27.6 Paa (0.004 psia); and

d) reducing at least some of the chlorine-complexed metal in the catalyst to the metallic state by contacting the chlorinated catalyst from step (c) with a gaseous stream comprising inert gas and hydrogen under reducing conditions at a total pressure below 690 kPaa (100 psia) and a temperature of from 350°C to 550°C to result in a regenerated catalyst comprising dispersed metal and zeolite.

2. A process according to claim 1 wherein step (a) is performed at a temperature of from 520°C to 600°C and/or at an oxygen partial pressure not greater than 69 kPaa (10 psia) and/or at a water partial pressure not greater than 10.3 kPaa (1.5 psia) and/or at a total pressure of from sub-atmospheric to 2.07 MPaa (300 psia) and/or for a time of from 10 to 100 hours.

3. A process according to claim 1 or 2 wherein step (a) is performed until the gaseous stream after contact with the catalyst has a carbon dioxide partial pressure below 69 Paa (0.01 psia), preferably below 27.6 Paa (0.004 psia).

4. A process according to any preceding claim in which step (a) is performed in a first stage (a1) and a second stage (a2) wherein stage (a1) is performed at a lower temperature than stage (a2).

5. A process according to claim 4 wherein (a1) is performed at a temperature of from 400°C to 500°C and (a2) is performed at a temperature of from 520°C to 600°C.
6. A process according to any preceding claim wherein step (a) is performed for a time of from 48 to 75 hours.

7. A process according to any preceding claim wherein step (b) is performed at a temperature of from 480° to 550°C and/or at an oxygen partial pressure not greater than 69 kPaa (10 psia) and/or at a water partial pressure not greater than 10.3 kPaa (1.5 psia) and/or at a total pressure of from sub-atmospheric to 2.07 MPaa (300 psia) and/or for at least 2 hours after HCl and/or Cl₂ breakthrough.

8. A process according to claim 7 wherein step (b) is performed at an oxygen partial pressure of from 13.8 to 27.6 kPaa (2 to 4 psia) and/or at a water partial pressure of from 1.38 to 5.17 kPaa (0.2 to 0.75 psia) and/or at a total pressure of from 345 kPaa to 1.38 MPaa (50 to 200 psia).

9. A process according to any preceding claim wherein step (b) is performed at a chlorine partial pressure of from 207 Paa to 20.7 kPaa (0.03 to 3 psia), preferably from 345 Paa to 6.9 kPaa (0.05 to 1 psia).

10. A process according to any preceding claim wherein step (c) is performed at a temperature of from 480° to 520°C and/or at a water partial pressure not greater than 10.3 kPaa (1.5 psia) and/or at an oxygen partial pressure of less than 31 kPaa (4.5 psia).

11. A process according to claim 10 wherein step (c) is performed at a water partial pressure of from 345 Paa to 2.76 kPaa (0.05 to 0.4 psia) and/or at an oxygen partial pressure of from 4.82 Paa to 20.7 kPaa (0.7 to 3 psia).

12. A process according to any preceding claim wherein step (c) is performed at a total pressure of from sub-atmospheric to 448 kPaa (65 psia), preferably at about atmospheric pressure.

13. A process according to any preceding claim wherein step (c) is performed until the hydrogen chloride in the gaseous stream after contact with the catalyst falls to a partial pressure below 10.3 Paa (0.0015 psia).

14. A process according to any preceding claim wherein step (d) is performed at a hydrogen partial pressure greater than 103 Paa (0.015 psia) and/or at a temperature of from 480° to 520°C and/or in the presence of water, preferably at partial pressure not greater than 10.3 kPaa (1.5 psia).

15. A process according to claim 14 wherein step (d) is performed at a hydrogen partial pressure of from 1.03 kPaa to 34.5 kPaa (0.15 to 5 psia) and/or at a water partial pressure of from 345 Paa to 2.76 kPaa (0.05 to 0.4 psia).

16. A process according to any preceding claim wherein step (d) is performed at a total pressure of from sub-atmospheric to 448 kPaa (65 psia), preferably at about atmospheric pressure.

17. A process according to any preceding claim wherein step (d) is performed until the hydrogen chloride in the gaseous stream after contact with the catalyst falls to a partial pressure below 51.7 Paa (0.0075 psia).

18. A process according to any preceding claim wherein between steps (a) and (b) there is performed an intermediate reducing step (a') comprising contacting the substantially decoked catalyst with a gaseous stream comprising inert gas and hydrogen under reducing conditions to reduce the metal component of the catalyst to the metallic state.

19. A process according to any preceding claim wherein between steps (c) and (d) there is performed a purging step (c') comprising contacting the chlorinated catalyst from step (c) with an oxygen-free displacement gas flow to purge oxygen therefrom prior to performing step (d).

20. A process according to any preceding claim wherein the catalyst comprises a type L zeolite optionally containing potassium or barium as exchangeable cations and/or the Group VIII catalytic metal comprises platinum and/or the Group VIII metal is present as from 0.05 to 6 wt% of the catalyst, which optionally comprises a binder.

21. A process for reforming hydrocarbons comprising contacting hydrogen and naphtha with a catalyst regenerated in accordance with the process according to any preceding claim, to produce aromatic hydrocarbons.

Patentansprüche

1. Verfahren zum Regenerieren eines desaktivierten Katalysators, der ein katalytisches Gruppe VIII Metall und Zeolith umfaßt, bei dem
a) Koks von dem desaktivierten Katalysator abgebrannt und das Metall in agglomerierte Teilchen überführt wird, die in der nachfolgenden Verfahrensstufe b) für chlorhaltiges Gas erreichbar sind, indem der Katalysator mindestens 6 Stunden mit einem gasförmigen Strom, der Sauerstoff, Inertgas und Wasser umfaßt, unter Oxi-
dationsbedingungen, die eine Temperatur von 400°C bis 600°C umfassen, kontaktiert wird, um den Katalysa
tor im wesentlichen zu entkoken und den Katalysator soweit zu agglomerieren, daß mindestens 80 Gew.% des
Metalls außerhalb der Kanäle des Zeolithen als Teilchen von mehr als 200 Å Größe agglomeriert sind,
b) das Metall des im wesentlichen entkokten Katalysators chloriert und dispergiert wird, indem der Katalysator
mit einem gasförmigen Strom, der Wasser, eine Chlorquelle, Sauerstoff und ein Inertgas umfaßt, unter Oxy-
chlorierungsbedingungen kontaktiert wird, die eine Temperatur von 450°C bis 550°C und einen Partialdruck
des Chlors, das aus der Chlorquelle stammt, von mindestens 207 Paa (0,03 psia) umfassen, bis der Chlora
erstoff und/oder das Chlor, der bzw. das hinter dem Katalysator durchbricht, einen Partialdruck von HCl + Cl₂
von mehr als 138 Paa (0,02 psia) erreichen, um das Metall im wesentlichen vollständig in den Chlorkomplex
to überführen und zu dispergieren,
c) mindestens ein Teil des Chlors von dem Metall entfernt wird, indem der chlorierte Katalysator mit einem gas-
förmigen Strom, der Wasser, Sauerstoff und ein Inertgas umfaßt, unter Chlorentfernungsbedingungen bei
einer Temperatur von 450°C bis 550°C und einem Gesamtdruck unter 690 kPaa (100 psia) kontaktiert wird, bis
der Chlorwasserstoff in dem gasförmigen Strom nach Kontakt mit dem Katalysator auf einen Partialdruck unter
27,6 Paa (0,004 psia) abfällt, und
d) mindestens ein Teil des als Chlorkomplex vorliegenden Metalls in dem Katalysator zum metallischen
Zustand reduziert wird, indem der chlorierte Katalysator aus Stufe (c) mit einem gasförmigen Strom, der Inert-
gas und Wasserstoff umfaßt, unter reduzierenden Bedingungen bei einem Gesamtdruck unter 690 kPaa (100
psia) und einer Temperatur von 350°C bis 550°C kontaktiert wird, um zu einem regenerierten Katalysator zu
führen, der dispergiertes Metall und Zeolith umfaßt.

2. Verfahren nach Anspruch 1, bei dem Stufe (a) bei einer Temperatur von 520°C bis 600°C und/oder einem Sauer-
stoff-Partialdruck von nicht mehr als 69 kPaa (10 psia) und/oder einem Wasser-Partialdruck von nicht mehr als
10,3 kPaa (1,5 psia) und/oder einem Gesamtdruck von unter atmosphärischem Druck liegendem Druck bis 2,07
MPa (300 psia) und/oder für einen Zeitraum von 10 bis 100 Stunden durchgeführt wird.

3. Verfahren nach Anspruch 1 oder 2, bei dem Stufe (a) durchgeführt wird, bis der gasförmige Strom nach Kontakt mit
dem Katalysator einen Kohlendioxid-Partialdruck unter 69 Paa (0,01 psia), vorzugsweise unter 27,6 Paa (0,004
psia) aufweist.

4. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (a) in einer ersten Stufe (a1) und einer zwei-
ten Stufe (a2) durchgeführt wird, wobei Stufe (a1) bei einer niedrigeren Temperatur als Stufe (a2) durchgeführt
wird.

5. Verfahren nach Anspruch 4, bei der (a1) bei einer Temperatur von 400°C bis 500°C durchgeführt wird und (a2) bei
einer Temperatur von 520°C bis 600°C durchgeführt wird.

6. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (a) für eine Zeitdauer von 48 bis 75 Stunden
durchgeführt wird.

7. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (b) bei einer Temperatur von 480°C bis
550°C und/oder bei einem Sauerstoff-Partialdruck von nicht größer als 69 kPaa (10 psia) und/oder bei einem Was-
er-Partialdruck von nicht größer als 10,3 kPaa (1,5 psia) und/oder bei einem Gesamtdruck von unter atmosphäri-
schem Druck liegendem Druck bis 2,07 MPa (300 psia) und/oder mindestens 2 Stunden nach dem Durchbruch
von HCl und/oder Cl₂ durchgeführt wird.

8. Verfahren nach Anspruch 7, bei dem Stufe (b) bei einem Sauerstoff-Partialdruck von 13,8 bis 27,6 kPaa (2 bis 4
psia) und/oder bei einem Wasser-Partialdruck von 1,38 bis 5,17 kPaa (0,2 bis 0,75 psia) und/oder einem Gesamt-
druck von 345 kPaa bis 1,38 MPa (50 bis 200 psia) durchgeführt wird.

9. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (b) bei einem Chlor-Partialdruck von 207
Paa bis 20,7 kPaa (0,03 bis 3 psia), vorzugsweise 345 Paa bis 6,9 kPaa (0,05 bis 1 psia) durchgeführt wird.

10. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (c) bei einer Temperatur von 480°C bis 520°C
und/oder einem Wasser-Partialdruck von nicht größer als 10,3 kPaa (1,5 psia) und/oder einem Sauerstoff-Partial-
druck von weniger als 31 kPaa (4,5 psia) durchgeführt wird.
11. Verfahren nach Anspruch 10, bei dem Stufe (c) bei einem Wasser-Partialdruck von 345 Paa bis 2,76 kPaa (0,05 bis 0,4 psia) und/oder einem Sauerstoff-Partialdruck von 4,82 Paa bis 20,7 kPaa (0,7 bis 3 psia) durchgeführt wird.

12. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (c) bei einem Gesamtdruck von unter atmosphärischem Druck liegendem Druck bis 448 kPaa (65 psia), vorzugsweise bei etwa atmosphärischem Druck durchgeführt wird.

13. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (c) durchgeführt wird, bis der Chlorwasserstoff in dem gasförmigen Strom nach Kontakt mit dem Katalysator auf einen Partialdruck unter 10,3 Paa (0,0015 psia) abfällt.

14. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (d) bei einem Wasserstoff-Partialdruck von mehr als 1,03 kPaa (0,015 psia) und/oder bei einer Temperatur von 480° bis 520°C und/oder in Gegenwart von Wasser, vorzugsweise bei einem Partialdruck von nicht mehr als 1,03 kPaa (1,5 psia) durchgeführt wird.

15. Verfahren nach Anspruch 14, bei dem Stufe (d) bei einem Wasserstoff-Partialdruck von 1,03 kPaa bis 34,5 kPaa (0,015 psia bis 5 psia) und/oder bei einem Wasser-Partialdruck von 345 Paa bis 2,76 kPaa (0,05 bis 0,4 psia) durchgeführt wird.

16. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (d) bei einem Gesamtdruck von unter atmosphärischem Druck liegendem Druck bis 448 kPaa (65 psia) und/oder bei einem Wasser-Partialdruck von 345 Paa bis 2,76 kPaa (0,05 bis 0,4 psia) durchgeführt wird.

17. Verfahren nach einem der vorhergehenden Ansprüche, bei dem Stufe (d) durchgeführt wird, bis der Chlorwasserstoff in dem gasförmigen Strom nach Kontakt mit dem Katalysator auf einen Partialdruck unter 51,7 Paa (0,0075 psia) abfällt.

18. Verfahren nach einem der vorhergehenden Ansprüche, bei dem zwischen den Stufen (a) und (b) eine Zwischendehaktionsstufe (a') durchgeführt wird, in der der im wesentlichen entkochte Katalysator mit einem gasförmigen Strom, der Inertgas und Wasserstoff umfaßt, unter reduzierenden Bedingungen für eine Zeitdauer kontaktiert wird, um die Metallokomponenten des Katalysators zum metallischen Zustand zu reduzieren.

19. Verfahren nach einem der vorhergehenden Ansprüche, bei dem zwischen den Stufen (c) und (d) eine Spülstufe (c') durchgeführt wird, bei der der chlorierte Katalysator aus Stufe (c) mit einer sauerstoffreien Verdrängungsströmung kontaktiert wird, um aus diesem Sauerstoff auszuspülen, bevor Stufe (d) durchgeführt wird.


Revendications

1. Procédé pour régénérer un catalyseur désactivé comprenant un métal catalytique du Groupe VIII et une zéolite, procédé qui comprend les étapes consistant :

a) à éliminer du catalyseur désactivé le coke par combustion et à transformer le métal en particules agglomérées qui sont accessibles à un gaz contenant du chlore dans l’étape b) suivante du procédé en mettant en contact le catalyseur pendant au moins 6 heures avec un courant gazeux comprenant de l’oxygène, un gaz inert et de l’eau dans des conditions d’oxydation comprenant une température de 400°C à 600°C pour provoquer une forte décokefaction du catalyseur et l’agglomération du catalyseur à un degré tel qu’au moins 80 % en poids du métal soient agglomérés à l’extérieur des canaux de la zéolite sous forme de particules ayant des dimensions supérieures à 200 Ångströms ;

b) à chlorer et disperser le métal du catalyseur ayant subi une forte décokefaction en mettant en contact le catalyseur avec un courant gazeux comprenant de l’eau, une source de chlore, de l’oxygène et un gaz inerte
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dans des conditions d’oxychloration comprenant une température de 450°C à 550°C et une pression partielle du chlore provenant de la source de chlore qui est égale à au moins 207 Paa (0,03 psia), jusqu’à ce que le chlorure d’hydrogène et/ou le chlore passant à travers le catalyseur atteignent une pression partielle de HCl + Cl₂ supérieure à 138 Paa (0,02 psia) pour parvenir pratiquement totalement à complexer avec le chlore et disperser le métal ;
c) à éliminer au moins une certaine quantité du chlore du métal en mettant en contact le catalyseur chloré avec un courant gazeux comprenant de l’eau, de l’oxygène et un gaz inerte dans des conditions d’élimination du chlore à une température de 450°C à 550°C et une pression totale inférieure à 690 kPaa (10 psia), jusqu’à ce que la pression du chlorure d’hydrogène dans le courant gazeux après la mise en contact avec le catalyseur tombe à une pression partielle inférieure à 27,6 Paa (0,004 psia) ; et
d) à réduire au moins une certaine quantité du métal complexé avec le chlore dans le catalyseur à l’état métallique en mettant en contact le catalyseur chloré de l’étape c) avec le courant gazeux comprenant un gaz inerte et de l’hydrogène dans des conditions réductrices à une pression totale inférieure à 690 kPaa (100 psia) et une température de 350°C à 550°C pour qu’il en résulte un catalyseur régnéré comprenant le métal dispersé et la zéolite.

2. Procédé suivant la revendication 1, dans lequel l’étape (a) est mise en œuvre à une température de 520°C à 600°C et/ou à une pression partielle d’oxygène non supérieure à 69 kPa (10 psia) et/ou à une pression partielle d’eau non supérieure à 10,3 kPa (1,5 psia) et/ou à une pression totale comprise dans l’intervalle d’une valeur inférieure à la pression atmosphérique à une valeur de 2,07 MPa (300 psia) et/ou pendant un temps de 10 à 100 heures.

3. Procédé suivant la revendication 1 ou 2, dans lequel l’étape (a) est mise en œuvre jusqu’à ce que le courant gazeux après la mise en contact avec le catalyseur ait une pression partielle de dioxyde de carbone inférieure à 69 Paa (0,01 psia), de préférence inférieure à 27,6 Paa (0,004 psia).

4. Procédé suivant l’une quelconque des revendications précédentes, dans lequel l’étape (a) est mise en œuvre dans une première étape (a1) et une seconde étape (a2), l’étape (a1) étant mise en œuvre à une température inférieure à celle de l’étape (a2).

5. Procédé suivant la revendication 4, dans lequel l’étape (a1) est mise en œuvre à une température de 400°C à 500°C et l’étape (a2) est mise en œuvre à une température de 520°C à 600°C.

6. Procédé suivant l’une quelconque des revendications précédentes, dans lequel l’étape (a) est mise en œuvre pendant un temps de 48 à 75 heures.

7. Procédé suivant l’une quelconque des revendications précédentes, dans lequel l’étape (b) est mise en œuvre à une température de 4800 à 550°C et/ou à une pression partielle d’oxygène non supérieure à 69 kPa (10 psia) et/ou à une pression partielle d’eau non supérieure à 10,3 kPa (1,5 psia) et/ou à une pression totale comprise dans l’intervalle d’une valeur inférieure à la pression atmosphérique à une valeur de 2,07 MPa (300 psia) et/ou pendant un temps d’au moins 2 heures après la percée de HCl et/ou de Cl₂.

8. Procédé suivant la revendication 7, dans lequel l’étape (b) est mise en œuvre à une pression partielle d’oxygène de 13,8 à 27,6 kPa (2 à 4 psia) et/ou à une pression partielle d’eau de 1,38 à 5,17 kPa (0,2 à 0,75 psia) et/ou à une pression totale de 345 kPa à 1,38 MPa (50 à 200 psia).

9. Procédé suivant l’une quelconque des revendications précédentes, dans lequel l’étape (b) est mise en œuvre à une pression partielle de chlore de 207 Paa à 20,7 kPa (0,03 à 3 psia), de préférence de 345 Paa à 6,9 kPa (0,05 à 1 psia).

10. Procédé suivant l’une quelconque des revendications précédentes, dans lequel l’étape (c) est mise en œuvre à une température de 480° à 520°C et/ou à une pression partielle d’eau non supérieure à 10,3 kPa (1,5 psia) et/ou à une pression partielle d’oxygène inférieure à 31 kPa (4,5 psia).

11. Procédé suivant la revendication 10, dans lequel l’étape (c) est mise en œuvre à une pression partielle d’eau de 345 Paa à 2,76 kPa (0,05 à 0,4 psia) et/ou à une pression partielle d’oxygène de 4,82 Paa à 20,7 kPa (0,7 à 3 psia).

12. Procédé suivant l’une quelconque des revendications précédentes, dans lequel l’étape (c) est mise en œuvre à une pression totale comprise dans l’intervalle d’une valeur inférieure à la pression atmosphérique à une valeur de
13. Procédé suivant l'une quelconque des revendications précédentes, dans lequel l'étape (c) est mise en oeuvre jusqu'à ce que la pression du chlorure d'hydrogène dans le courant gazeux après la mise en contact avec le catalyseur tombe à une pression partielle inférieure à 10,3 Paa (0,015 psia).

14. Procédé suivant l'une quelconque des revendications précédentes, dans lequel l'étape (d) est mise en oeuvre à une pression partielle d'hydrogène supérieure 103 Paa (0,015 psia) et/ou à une température de 480° à 520°C et/ou en présence d'eau, de préférence à une pression partielle non supérieure à 10,3 kPaa (1,5 psia).

15. Procédé suivant la revendication 14, dans lequel l'étape (d) est mise en oeuvre à une pression partielle d'hydrogène de 1,03 kPaa à 34,5 kPaa (0,15 à 5 psia) et/ou à une pression partielle d'eau de 345 Paa à 2,76 kPaa (0,05 à 0,4 psia).

16. Procédé suivant l'une quelconque des revendications précédentes, dans lequel l'étape (d) est mise en oeuvre à une pression totale comprise dans l'intervalle d'une valeur inférieure à la pression atmosphérique à une valeur de 448 kPaa (65 psia), de préférence approximativement égale à la pression atmosphérique.

17. Procédé suivant l'une quelconque des revendications précédentes, dans l'étape (d) est mise en oeuvre jusqu'à ce que la pression du chlorure d'hydrogène dans le courant gazeux après la mise en contact avec le catalyseur tombe à une pression partielle inférieure à 51,5 Paa (0,0075 psia).

18. Procédé suivant l'une quelconque des revendications précédentes, dans lequel, entre les étapes (a) et (b), on met en oeuvre une étape de réduction intermédiaire (a') comprenant la mise en contact du catalyseur ayant subi une forte décokefaction avec un courant gazeux comprenant un gaz inerte et de l'hydrogène dans des conditions réductrices pour réduire le constituant métallique du catalyseur à l'état métallique;

19. Procédé suivant l'une quelconque des revendications précédentes, dans lequel, entre les étapes (c) et (d), on met en oeuvre une étape de purge (c') comprenant la mise en contact du catalyseur chloré de l'étape (c) avec un courant gazeux de déplacement dépourvu d'oxygène pour chasser l'oxygène de ce catalyseur avant la mise en oeuvre de l'étape (d).

20. Procédé suivant l'une quelconque des revendications précédentes, dans lequel le catalyseur comprend une zéolite de type L contenant facultativement du potassium ou du baryum comme cations échangeables et/ou le métal catalytique du Groupe VIII comprend le platine et/ou le métal du Groupe VIII est présent en une teneur de 0,05 à 6 % en poids du catalyseur, qui comprend facultativement un liant.

21. Procédé pour le reformage d'hydrocarbures, comprenant la mise en contact d'hydrogène et d'un naphta avec un catalyseur régénéré conformément au procédé suivant l'une quelconque des revendications précédentes, pour produire des hydrocarbures aromatiques.