



(11) **EP 1 456 154 B9**

(12) **CORRECTED EUROPEAN PATENT SPECIFICATION**

(15) Correction information:
Corrected version no 1 (W1 B1)
Corrections, see
Description Paragraph(s) 24, 88, 96, 111

(51) Int Cl.:
C07C 2/86 ^(2006.01) **B01J 29/072** ^(2006.01)
B01J 29/06 ^(2006.01)

(48) Corrigendum issued on:
15.10.2014 Bulletin 2014/42

(86) International application number:
PCT/EP2002/014817

(45) Date of publication and mention
of the grant of the patent:
23.04.2014 Bulletin 2014/17

(87) International publication number:
WO 2003/053892 (03.07.2003 Gazette 2003/27)

(21) Application number: **02796749.6**

(22) Date of filing: **18.12.2002**

(54) **PROCESS FOR THE ALKYLATION OF BENZENE**

VERFAHREN ZUR ALKYLIERUNG VON BENZEN

PROCEDE D'ALKYLATION DE BENZÈNE

(84) Designated Contracting States:
AT BE BG CH CY CZ DE DK EE ES FI FR GB GR
IE IT LI LU MC NL PT SE SI SK TR

(30) Priority: **20.12.2001 IT MI20012707**

(43) Date of publication of application:
15.09.2004 Bulletin 2004/38

(73) Proprietor: **versalis S.p.A.**
20097 San Donato Milanese (MI) (IT)

(72) Inventors:
• **GIROTTI, Gianni**
I-28100 Novara (IT)
• **RIVETTI, Franco**
I-20139 Milan (IT)
• **RAMELLO, Stefano**
I-28100 Novara (IT)

(74) Representative: **Bottero, Carlo et al**
Barzanò & Zanardo Milano S.p.A.
Via Borgonuovo, 10
20121 Milano (IT)

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Description

[0001] The present invention relates to a process for the alkylation of benzene which comprises reacting benzene with acetone and hydrogen in the presence of a catalytic composition comprising a solid acid material and copper. A preferred aspect is for the catalytic composition to also contain one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides. A particularly preferred aspect is for the catalytic composition to contain one or more elements selected from elements of groups IIIA and VIB.

[0002] In particular, the invention relates to a process for the production of cumene starting from the reagents acetone, benzene and hydrogen, which are reacted in a single reaction step in the presence of said catalytic system.

[0003] Even more specifically, the invention relates to an alkylation process of benzene, with acetone and hydrogen in the presence of a catalytic composition containing a solid acid material, copper and one or more elements selected from Cr and Al, wherein the solid acid material comprises or consists of a zeolite, preferably zeolite beta.

[0004] The new preparation of cumene according to the present invention can be used in particular in a production process of phenol comprising the following steps:

(a) reacting benzene, acetone and hydrogen in the presence of the catalytic system according to what is specified above, comprising a solid acid material and copper,

(b) oxidizing the cumene to cumene hydroperoxide,

(c) treating the cumene hydroperoxide with acids to obtain a re-arrangement to phenol and acetone.

[0005] A preferred aspect is for the catalytic composition used in step (a) to also contain one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides. A particularly preferred aspect is for the catalytic composition of step (a) to contain one or more elements selected from elements of groups IIIA and VIB.

[0006] The acetone which is formed in step (c) can be recycled to step (a) for the synthesis of cumene.

[0007] According to what is specified above, the catalytic composition used in step (a) preferably comprises zeolite beta and copper. A particularly preferred aspect is to use in step (a) a catalytic composition containing zeolite beta, Cu and a metal selected from Cr and Al.

[0008] Cumene or isopropyl benzene is an important intermediate in the basic chemical industry mainly used as precursor for the production of phenol, in turn useful as intermediate in the preparation of caprolactam from which nylon is produced.

[0009] The industrial synthesis of phenol comprises the alkylation steps of benzene to cumene, the oxidation of cumene to cumyl hydroperoxide and the subsequent rearrangement to give phenol and acetone.

[0010] As far as the alkylation of benzene to cumene is concerned, catalysts based on phosphoric acid and infusorial earth for fixed bed reactors or $AlCl_3$ in slurry, and propylene as alkylating agent, are still widely used in the petrochemical industry.

[0011] These processes however create problems relating to environmental impact and safety: the use of these catalysts, in fact, is particularly problematical due to corrosion, the by-production of toxic organic products and the disposal of the exhausted catalysts.

[0012] In 1965, however, the preparation of cumene was described for the first time, using zeolite X or zeolite Y as catalyst (Minachev, Kr. M., et al, Neftekhimiya 5 (1965) 676). The use of zeolites with a faujasitic structure for the alkylation of benzene with light olefins such as propylene, was subsequently described by Venuto et al. (J. Catal. 5, (1966) 81).

[0013] U.S. 4,292,457 describes the use of ZSM-5 type zeolites for alkylating benzene with propylene.

[0014] US-A-2,410,553 discloses a process for preparing alkylated aromatic compounds by reacting an aromatic compound and a ketone in presence of a catalyst, in particular of a zinc chloride catalyst.

[0015] US-A-2,412,230 discloses a process for producing aromatic hydrocarbons subjecting an aromatic hydrocarbon into contact with an alkoxy compound under alkylating condition in presence of a catalytically effective amount of pyrophosphate of a heavy metal selected from the members of the right-hand column of Group I of the Periodic Table.

[0016] Excellent results in terms of industrial application have been obtained in the synthesis of cumene using zeolites with a beta-type structure, as described in EP 432814, and in particular using catalysts comprising beta zeolite according to what is described in EP 687500 and in EP 847802.

[0017] Once cumene has been obtained, it is transformed into phenol by means of an oxidation step to cumene hydroperoxide, followed by an acid treatment step which causes the breakage of the peroxide bond with the formation of phenol and acetone.

[0018] The synthesis of phenol via cumene, on which most industrial plants existing throughout the world for the production of phenol, are based, leads to the coproduction of a quantity of acetone equal to 0.61 kg per kg of phenol.

[0019] Phenol is mainly used in the production of bisphenol A (about 35%), phenolic resins (about 35%), caprolactam (about 15%), aniline, alkylphenols, xlenols and other products, whereas acetone is mainly used in the production of methylmethacrylate (about 45%), bisphenol A (about 20%), solvents (about 17%) and methylisobutylketone (about 8%).

[0020] There is therefore an unbalanced situation, at least potential, in the request for phenol and acetone, intrinsically deriving from their co-production, which does not allow a modulation in their supply in relation to the growth margins of the different sectors and outlet markets for the two products.

[0021] New processes based on the re-use of acetone - co-produced with phenol - in the upstream synthesis of cumene, have been proposed to avoid this situation. In EP 361755, the propylene used as alkylation agent of benzene for the synthesis of cumene is obtained, either totally or partially, starting from acetone, after reduction with hydrogen to isopropanol and subsequent dehydration to propylene.

[0022] The re-use of the possible excess acetone for the re-production of propylene according to the method described above, is extremely onerous, particularly due to the high number of steps associated with the chemical reduction transformations to isopropyl alcohol and the subsequent dehydration of the alcohol to propylene.

[0023] An alternative which reduces the number of chemical transformations necessary for the re-use of acetone consists in the direct use of isopropanol, obtained by the reduction of acetone with hydrogen, as alkylation agent of benzene in the synthesis of cumene, as described for example in EP 1069100.

[0024] WO01/62692 relates to a process for the preparation of cumene by reacting isopropanol or a mixture of isopropanol and propene with benzene in the presence of a beta-zeolite catalyst having a SiO₂/Al₂O₃ molar ratio greater than 10:1. Also a process is taught for preparing phenol from benzene, which comprises the steps:

- I. Preparation of cumene according to the above process,
- II. Oxidation of cumene to cumene hydroperoxide,
- III. acid-catalyzed cleavage of cumene hydroperoxide to give phenol and acetone and
- IV. hydrogenation of acetone to form isopropanol.

[0025] EP1069100 relates to a process for the alkylation of aromatic compounds by the reaction of the aromatic compound of interest with isopropanol, alone or mixed with propylene, consisting in carrying out said reaction in the presence of a catalytic composition based on zeolite, under mixed gas-liquid phase conditions or under completely liquid phase conditions, at such temperature and pressure that the concentration of water in the reaction liquid phase is not higher than 8,000 ppm, regardless of the total water content present in the reaction mixture. A process for the preparation of phenol is also taught comprising said alkylation process, the oxidation of the cumene thus obtained, the acid treatment of cumylhydroperoxide, the hydrogenation of the acetone which is formed as by-product and recycling of the isopropanol which is thus formed.

[0026] The direct alkylation of benzene with isopropanol, obtained by the reduction of acetone co-produced with phenol, represents an improvement, from an industrial point of view, with respect to the option which involves the re-production of propylene to be used as alkylation agent of benzene, but the best solution, from an industrial and process point of view, would obviously consist of the direct use of acetone as alkylation agent of benzene, in the presence of hydrogen, in the synthesis of cumene.

[0027] A catalytic system has now been found, which is capable of promoting the synthesis of cumene starting directly from the reagents acetone, benzene and hydrogen.

[0028] The cumene, obtained according to the industrial process claimed herein, can be used for the subsequent production of phenol by means of oxidation to cumene hydroperoxide and the subsequent rearrangement of the hydroperoxide to phenol and acetone. The acetone thus obtained, can in turn be used for the synthesis of cumene by the direct alkylation of benzene with acetone and hydrogen carried out according to the industrial process claimed herein.

[0029] Therefore, the object of the present invention relates to a process for the alkylation of benzene to give cumene which comprises reacting acetone, benzene and hydrogen, which are reacted in a single reaction step, in the presence of a catalytic system comprising a solid acid material and copper. A preferred aspect is for the catalytic composition used for the alkylation of benzene to cumene to additionally contain one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides. A particularly preferred aspect is for the catalytic composition to contain one or more elements selected from elements of groups IIIA and VIB, preferably Cr or Al.

[0030] The alkylation process, object of the present invention, allows the preparation of phenol from cumene to be considerably simplified; it is not necessary in fact to first effect the chemical reduction transformation of acetone to isopropanol as described for example in U.S. 5,160,497, the subsequent chemical dehydration transformation of isopropanol to propylene as described for example in U.S. 5,017,729 and consequently the subsequent chemical alkylation transformation of benzene with propylene as described for example in EP 439632.

[0031] The chemical transformations described above are normally carried out under somewhat different reaction conditions for each of the reactions in question and in the presence of equally different catalysts.

[0032] The catalytic system used in the process of the present invention allows all the chemical transformations necessary for the preparation of cumene starting from the reagents acetone, benzene and hydrogen, to be contemporaneously carried out in a single reaction step, maximizing the yield to cumene and minimizing the secondary reactions of the various reagents, intermediates and products.

[0033] Particularly significant secondary and undesired reactions for determining the overall yield which could be expected by experts in the field are the parallel reduction reaction of benzene with hydrogen to cyclohexene, cyclohexane and hexane, the parallel condensation reaction of acetone to 4-methyl-3-penten-2-one and the subsequent reactions of these by-products with the various reagents and products of the main reactions such as for example the alkylation of benzene to phenylcyclohexane due to the cyclohexene and reduction of 4-methyl-3-penten-2-one to 4-methyl-2-pentanone and 4-methyl-2-pentanol due to the hydrogen.

[0034] The catalytic system used in the process of the present invention unexpectedly allows the conversion of the reagents to be oriented towards the desired product, reducing the formation of undesired products to the minimum.

[0035] The catalytic system used in the alkylation process of benzene, which is object of the present invention, comprises a solid acid material and copper, wherein the copper is preferably in the form of an oxide.

[0036] According to a preferred aspect, the catalytic composition also contains one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides. These elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, are also preferably in the form of oxides. In particular, copper and these elements can be contained in the catalytic composition in the form of a mixed oxide.

[0037] According to a particularly preferred aspect, the catalytic composition contains one or more elements selected from the elements of groups IIIA and VIB. In accordance with what is specified above, these elements of groups IIIA and VIB are preferably in the form of oxides. In particular, copper and these elements can be contained in the catalytic composition in the form of a mixed oxide.

[0038] A particularly preferred aspect of the present invention is to use catalytic compositions containing copper and an element selected from chromium and aluminum. In particular, copper and these elements can be contained in the catalytic composition in the form of a mixed oxide.

[0039] In particular, copper and chromium can be contained in the catalytic composition in the form of copper chromite. Copper chromite is represented by the empirical formula $\text{CuO} \cdot \text{CuCr}_2\text{O}_4$. CuCr_2O_4 is known as C.A.S. R.N. 12018-10-9 and is described in "Gmelins Handbuch der An-organischen Chemie, 8th ed., Vol. Kupfer, part B, Installment 3, system number 60, page 60". In the process of the present invention, commercially available materials called copper chromite, can be used, containing Cu (II) and Cr (III), having varying proportions of CuO and CuCr_2O_4 . These materials, which can optionally also contain small quantities of promoters such as Ba and Mn, are well known to experts in the field and are described for example in J.D. Stoupe, "An X-Ray Diffraction Study of the Copper Chromites and of the "Copper-Chromium Oxide" Catalyst" J.Am.Che.Soc., vol. 71, 1949, page 589; in A. Imura et al., "Catalysis by "Copper Chromite", I, The effect of hydrogen Reduction on the composition, structure and catalytic activity for methanol decomposition", Bull. Chem. Soc. Jp., 56, 2203-2207 (1983); in R.B.C. Pillai, "A study of the pre-activation of a copper chromite catalyst", Catalysis Letters 26 (1994) 365-371.

[0040] Copper and aluminum can be contained in the catalytic composition used in the present invention in the form of the corresponding oxides.

[0041] In accordance with what is specified above, the catalytic composition containing copper chromite can contain, as promoters, barium and/or manganese, preferably in the form of oxides. The barium or manganese content is lower than 15% by weight with respect to the total weight of the composition and preferably ranges from 0.1 to 5% by weight. The weight percentages of barium and manganese refer to their content expressed as element.

[0042] The solid acid material contained in the catalytic composition used in the alkylation process of benzene of the present invention, is preferably of a zeolitic nature and can contain one or more zeolitic materials. Zeolites which can be used are zeolite beta, zeolite Y, ZSM-12 and mordenite. These zeolites are described in "Atlas of zeolite structure types", Ch. Baerlocher, W.M.Meier and D.H. Olson, 2001, 5th Edition, Elsevier. A preferred aspect is to use zeolite beta.

[0043] The zeolites are used in acid form, i.e. in the form in which all the negative charges deriving from the aluminum present in the structure are counterbalanced by hydrogen ions, or prevalently acid.

[0044] The zeolite beta used as component of the catalytic composition of the process according to the present invention corresponds to that described in U.S. 3,308,069, and is a porous crystalline material having the composition



wherein n is the oxidation state of M, x is less than 1, y ranges from 5 to 100, w ranges from 0 to 4, M is a metal selected from those of groups IA, IIA, IIIA of the Periodic System or from transition metals and TEA is tetraethyl ammonium hydroxide.

[0045] A preferred aspect of the present invention is for the zeolite beta to be in acid form, i.e. in the form in which the

H⁺ ion has partially or totally substituted the metallic cations initially present.

[0046] This substitution is effected according to the known methods by means of an exchange with ammonium ions, washing and subsequent calcination.

[0047] The catalytic compositions which can be used in the alkylation process of the present invention, comprising a solid acid material and copper, can comprise suitable binding agents, for example oxides of groups IIIA, IVA and IVB. More preferably, the catalytic system can contain an oxide of Si or Al as binding carrier. Even more preferably, the catalytic system can contain γ -alumina as binding carrier.

[0048] γ -alumina is a known material and is commercially available in the form, preferred for the purposes of the invention, of the precursors bohemite or p-bohemite, transformed subsequently to γ -alumina during the preparation of the catalytic system, in the final calcination phase.

[0049] The binder is preferably used in a relative quantity by weight with respect to the catalytic system ranging from 5:95 to 95:5.

[0050] A particularly preferred aspect of the present invention is to use a catalytic system containing copper chromite and a beta-type zeolite, in its acid form. This composition can contain an inorganic binder in accordance with what is described above.

[0051] The copper is preferably contained in the catalytic composition of the present invention in a weight ratio of the metal with respect to the solid acid material ranging from 0.001 to 10, more preferably ranging from 0.01 to 2. When the catalytic composition contains one or more elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, each element is preferably contained in a weight ratio of the metal with respect to the solid acid material ranging from 0.001 to 10, more preferably from 0.01 to 2.

[0052] The catalytic system used in the present invention, as described above, contains a solid acid component with an alkylation functionality and a metallic component containing copper and optionally one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, with a hydrogenation functionality.

[0053] The catalytic system used in the present invention, containing a solid acid component and a metallic component containing copper and optionally one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, can be prepared, starting from the components described above, according to various practical combination procedures, each maintaining the specific characteristics listed above.

[0054] The catalytic system of the present invention can therefore consist of one or more distinct zones each containing a single functionality, either hydrogenation linked to the metallic component or alkylation linked to the acid component, in particular zeolitic, or both hydrogenation and alkylation functionalities having the characteristics described above.

[0055] Examples of some of the various preparation procedures of the catalytic system are indicated below and are schematically represented in figure 1.

[0056] In this figure, I refers to the metallic component of the catalytic composition, containing copper and optionally one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, said component having a hydrogenation functionality. When the metallic component contains copper alone, this is preferably in the form of an oxide. When the metallic component also contains an element selected from groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, it is also preferably in the form of an oxide. In this case, the component can be prepared for example by the mechanical mixing of the oxides. In the particular case in which the copper and the element or elements selected from groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, are in the form of a mixed oxide, component I can be prepared for example according to the known coprecipitation techniques, or by melting the metal oxides present in the mixed oxide. The metallic component I preferably consists of copper chromite.

[0057] Again with reference to figure 1, A refers to the solid acid component, preferably zeolitic, containing the alkylation functionality. Zeolites which can be used are preferably zeolite beta, zeolite Y, ZSM-12 and mordenite. A preferred aspect is to use zeolite beta. The solid acid component, preferably zeolitic, can be used in a mixture with suitable binding agents, for example oxides of the elements of groups IIIA, IVA and IVB. The solid acid component A more preferably contains an oxide of Si or Al as binding carrier. Even more preferably, the solid acid component A contains γ -alumina as binding carrier. The zeolitic composition with binder can be prepared according to any of the known techniques. In the case of zeolite beta, it can be prepared, for example, as described in EP 687500 and EP 847802.

[0058] Again with reference to figure 1, AI refers to a composition containing both hydrogenation and alkylation functionalities. The composition indicated with AI can also comprise suitable binding agents, for example oxides of groups IIIA, IVA and IVB. The composition indicated with AI more preferably contains an oxide of Si or Al as binding carrier. Even more preferably, the composition AI contains γ -alumina as binding carrier. The composition AI can be prepared according to any of the techniques well known to experts in the field, such as for example a) impregnation, b) ion exchange or c) extrusion, described hereunder:

a) it is possible to operate for example by impregnating the solid acid material, preferably of a zeolitic nature, with an aqueous solution containing a copper salt and optionally a salt of an element selected from groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, drying and calcining the resulting product. Separate solutions for the copper salt and salt of the element of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, can be used. The product obtained with this method can be optionally used in a mixture with a suitable binding agents, for examples oxides of groups IIIA, IVA and IVB, as described above. Alternatively, it is possible to operate by impregnating a mixture of solid acid material and binding agent with an aqueous solution containing a copper salt and optionally a salt selected from groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, drying and calcining the resulting product.

b) When the ion exchange technique is used, the solid acid material, preferably of a zeolitic nature, is for example put in an aqueous solution containing a copper salt and optionally a salt selected from groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, and the mixture left under stirring for a few hours. The solid in suspension is recovered by filtration, washed with demineralized water and dried: a solid acid material is obtained in exchanged form with copper ions and possibly ions of a metal selected from groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides. The material obtained in this way can optionally be used in a mixture with a suitable binding agent, as defined above. Alternatively, a composition of a solid acid material and of suitable binding agents is used, which is subjected to the ion exchange process described above.

c) Alternatively, an extrusion process can be used, in which a mechanical mixture of the two components, i.e. the solid acid material, preferably zeolitic, and the metallic component, is paste-mixed with a peptizing acid solution, extruded, dried and calcined with any traditional method. The product obtained in this way can optionally be used in a mixture with a suitable binding agent, defined above. Alternatively, a mechanical mixture of the three components, i.e. the solid acid material, preferably zeolitic, the metallic component and a suitable binding agent, defined above, is subjected to an extrusion process. In accordance with what is specified above, the catalytic system can consist for example (Figure 1a) of a sole Al composition zone containing both metallic and acid components, having alkylation and hydrogenation functionalities.

[0059] According to another embodiment, the catalytic system can consist for example (Figure 1b) of two or more separate Al composition zones, each of which contain both the acid and metallic components, having alkylation and hydrogenation functionalities, wherein the composition of the single zones differs in terms of chemical nature and proportion between the alkylation and hydrogenation functionality. Each single zone of the Al composition is prepared with one of the known methods described above, and the zones are then assembled by stratification inside the reactor.

[0060] Another preparation procedure (Figure 1c) of the catalytic system consists for example of two or more distinct zones, in one of which there is the metallic component I containing the catalytic hydrogenation function alone, whereas in the remaining zones there are one or more compositions Al with different combinations of the two components having alkylation and hydrogenation functionalities. Each single zone of the composition Al, containing two functionalities, is prepared with one of the known methods described above.

[0061] In another embodiment (Figure 1d), the catalytic system consists for example of two or more distinct zones, in one of which there is component I containing the catalytic hydrogenation function alone, whereas in the subsequent zone there is component A, having an alkylation catalytic function. The acid component A, preferably zeolitic, can be used in a mixture with suitable binding agents, defined above.

[0062] In a further embodiment (Figure 1e) the catalytic system consists for example of a single zone in which the two components of the catalytic composition, I and A respectively, mechanically mixed with each other, are arranged. Also in this case, the solid acid component A, preferably zeolitic, can be used in a mixture with suitable binding agents defined above.

[0063] In yet another embodiment (Figure 1f), the catalytic system consists for example of two or more distinct zones, in one of which there is component I of the catalytic system containing the catalytic hydrogenation function alone whereas in the remaining zones there are one or more different pairs of components I and A, mechanically mixed, each of which contains the catalytic hydrogenation function alone or the catalytic alkylation function alone. Also in this case, the solid acid component, preferably zeolitic, can be used in a mixture with suitable binding agents defined above.

[0064] When the catalytic system has a zone containing the metallic component I alone, having a catalytic hydrogenating function, this is preferably the one which first comes into contact with the stream of benzene, acetone and hydrogen. Furthermore, it is evident that, on the basis of the procedures described, a catalytic system characterized by the contemporaneous presence of a hydrogenation function, whose activity gradient decreases in one direction, and an alkylation function, whose activity gradient decreases in the opposite direction, can be easily produced by experts in the field, if desired.

[0065] The activity gradient of the hydrogenation function preferably decreases along the feeding direction and flow

of acetone, benzene and hydrogen, whereas the activity gradient of the alkylation function decreases along the opposite direction and flow.

[0066] According to a preferred aspect of the alkylation process of the present invention, it is preferable to operate at a reaction temperature generally ranging from 50 to 350°C, preferably from 100 to 250°C. The pressure is generally equal to or higher than atmospheric pressure and preferably ranges from 1 to 50 bars. A molar ratio between benzene and acetone, is used in the feeding, not lower than 1 and preferably higher than 2. A molar ratio between hydrogen and acetone, is used in the feeding not lower than 1 and preferably higher than 2. The catalytic composition is preferably pre-activated in a stream of hydrogen.

[0067] The reaction can be conveniently carried out in fixed bed catalyst reactors, containing one or more catalytic beds. The reagents can in this case all be fed to the reactor, in the desired proportions, to the first catalytic bed, or the feeding of the reagents or some of them can be partially fed to the different catalytic beds.

[0068] Some of the compositions prepared with the methods described above however can also be conveniently used in reactors different from fixed bed reactors.

[0069] According to the process claimed herein with reference to Figure 2, benzene, acetone and hydrogen are reacted in a single step in the presence of the catalytic system described and under the conditions indicated for obtaining a reaction product which mainly contains isopropylbenzene, non-converted benzene, non-converted hydrogen, water and polyisopropylbenzenes.

[0070] The reaction product is fractionated in a separation section S using conventional separation methods, such as degassing, distillation or the demixing of liquids, to obtain a first fraction mainly containing hydrogen, a second fraction mainly containing water, a third fraction mainly containing benzene, a fourth fraction mainly containing isopropylbenzene and a fifth fraction mainly containing polyisopropylbenzenes.

[0071] The first fraction (containing hydrogen) is re-used in the reaction step with acetone and benzene, the second fraction (containing water) is removed from the process, the third fraction (containing benzene) is partly re-used in the reaction step with acetone and hydrogen and partly in a subsequent reaction step, called transalkylation step, where it is reacted with the fifth fraction (containing polyisopropylbenzenes) to produce again the desired product isopropylbenzene.

[0072] Transalkylation is a reaction which is well known in the state of the art and is carried out in the presence of a solid acid catalyst, preferably in the presence of a solid acid catalyst based on zeolites, more preferably in the presence of a solid acid catalyst based on beta-type zeolite as described for example in EP 687500 and in EP 847802.

[0073] The temperature conditions for the transalkylation reaction are selected from 100 to 350°C, the pressure is selected from 10 to 50 atm and the WHSV ranges from 0.1 to 200 hours⁻¹, as also described in EP 687500 and in EP 847802.

[0074] The transalkylation reaction product is fractionated using the conventional separation methods in the same separation section.

[0075] The third fraction coming from the separation section therefore contains non-converted benzene coming from both the alkylation step and the transalkylation step. The fourth fraction coming from the separation section contains cumene coming from both the alkylation step and the transalkylation step and the fifth fraction coming from the separation step contains polyisopropylbenzenes coming from both the alkylation step and the transalkylation step.

[0076] The cumene obtained according to the process, object of the present invention, can be used for the production of phenol by means of oxidation to cumene hydroperoxide and the subsequent rearrangement of the hydroperoxide to phenol and acetone.

[0077] A further object of the present invention therefore relates to a process for the production of phenol which comprises the following steps:

- (a) reacting benzene, acetone and hydrogen in the presence of a catalytic system comprising a solid acid material and copper
- (b) oxidation of cumene to cumene hydroperoxide
- (c) rearrangement of the cumene hydroperoxide to phenol and acetone.

[0078] The acetone which is formed in step (c) can be recycled to the synthesis step (a) of cumene.

[0079] The catalytic composition used in step (a) also preferably contains one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, and is in accordance with what is specified above with respect to the alkylation process of the present invention. The catalytic composition used preferably contains zeolite beta, copper chromite and optionally an inorganic binder.

[0080] The various passages for the oxidation of cumene to cumene hydroperoxide, the rearrangement of hydroperoxide to give phenol and acetone and the purification of phenol are well known in literature, as described for example in U.S. 5,160,497 and U.S. 5,017,729.

[0081] The oxidation step of cumene to cumene hydroperoxide can be carried out for example with molecular oxygen

at a temperature ranging from 60 to 150°C and at a pressure ranging from 1 to 10 kg-f/cm². It is preferable to operate in the presence of an initiator and an alkaline compound for the pH control.

[0082] The transformation step of cumene hydroperoxide to phenol and acetone is carried out in the presence of an acid, for example a strong acid such as sulfuric acid, or for example an exchange resin or a silico-alumina. At the end, the reaction mixture is concentrated to recover the acetone, which can then be recycled to the alkylation step (a). It can be clearly seen how a process for the production of phenol starting from hydrogen, oxygen and benzene, with the coproduction of water alone, can actually be effected by recycling the acetone to step (a).

[0083] According to a preferred aspect, at the end of the first step, after separating the desired product, cumene, by fractionation, which passes to the subsequent oxidation step, the remaining fraction of polyisopropylbenzenes is used in a separate step for a transalkylation reaction with benzene to recover other cumene.

[0084] The transalkylation reaction is carried out in the presence of zeolite beta or a catalyst based on zeolite beta, in particular prepared according to what is described in EP 687500 and EP 847802, which also describe the reaction conditions.

[0085] The catalytic composition of the present invention containing a zeolite in acid form, preferably zeolite beta, Cu, and one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides with the proviso that when zeolite is zeolite Y and the catalytic composition contains one or more elements of Group VIB, said element is selected between Mo and W, is new and is a further aspect of the present invention.

[0086] This catalytic composition preferably contains zeolite beta, Cu and an element selected from Cr and Al. The metals contained in the composition are preferably in the form of oxides.. According to a particular aspect of the present invention, copper and chromium are contained in the form of copper chromite.

[0087] The copper is preferably contained in the catalytic composition of the present invention in a weight ratio of the metal with respect to the zeolite ranging from 0.001 to 10, more preferably ranging from 0.01 to 2. The catalytic composition contains one or more elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides with the proviso that when zeolite is zeolite Y and the catalytic composition contains one or more elements of Group VIB, said element is selected between Mo and W, each element is preferably contained in a weight ratio of the metal with respect to the zeolite ranging from 0.001 to 10, more preferably from 0.01 to 2. When the metallic component of the catalytic composition used in this patent comprises copper chromite, Ba and Mn can be present as promoters. These catalytic compositions may additionally contain a binder. The preparation of these catalytic compositions is in accordance with all the methods previously described.

[0088] The following examples are provided for a further illustration of the invention without limiting its scope in any way.

EXAMPLE 1 - Preparation of the catalytic system

[0089] A catalytic system is prepared, consisting of 10 g of a catalyst based on copper chromite (produced by the company SüdChemie with the trade-name G99b) having the following composition expressed as weight percentage of the elements: Cu 35%, Cr 31%, Ba 2%, Mn 2.5%, indicated hereunder as "material A1" and 4.5 g of a catalyst based on zeolite beta prepared according to the indications described in Example 1 of EP 687500, called "material B1". The zeolite beta used in the preparation of the material B1 is a product of the company Zeolyst with the trade-name CP-806 BL 25.

[0090] The catalytic system is prepared so that the first zone of said system consists of the material A1 alone in a quantity equal to 3 g and the second zone consists of a mechanical mixture of materials A1 and B1 in a quantity equal to 7 g of material A1 and 4.5 g of material B1. The total quantity of the catalytic system thus prepared is therefore equal to 14.5 g.

EXAMPLE 2 - Preparation of the catalytic system

[0091] A catalytic system is prepared, starting from the materials G-99b (57 g)) and zeolite beta (80 g), already used in the previous example, and alumina p-bohemite commercialized by the Company Laroche with the trade-name Versal 450 (143 g).

[0092] The materials are mechanically mixed in a ploughshare mixer for about 25 minutes, after which 175 cc of an aqueous solution of acetic acid at 5% w/w are added to the mixture of powders thus obtained, without interrupting the mixing for a further 20 minutes approximately.

[0093] The intermediate thus obtained is then subjected to extrusion using a HUTT type gear press extruder and the product thus obtained is subsequently subjected to aging for a time of not less than 48 hours.

[0094] After the period of aging, the product, in the form of pellets, is then subjected to calcination treatment in air at a temperature of 550°C approx. for 5 hours, obtaining a material called "material B2", containing about 25% of material G99b, about 30% of zeolite beta and about 45% of alumina binding carrier.

[0095] The catalytic system is prepared so as to consist of a single zone in which there is material B2. The total quantity of catalytic system thus prepared is equal to 9.0 g.

EXAMPLE 3 - Preparation of the catalytic system

[0096] A catalytic system is prepared, consisting of two distinct zones, the first of which contains 3 g of material A1 already used in Example 1 and the second containing 8 g of material B2 prepared according to the procedure described in Example 2. The total quantity of catalytic system is therefore equal to 11 g.

EXAMPLE 4 - Preparation of the catalytic system

[0097] 37 g of a catalyst based on zeolite beta prepared according to the indications provided in Example 1 of EP 687500 (12-16 mesh), are charged into a rotavapor flask having a volume of 500 ml, and are then dried under vacuum at room temperature for 2 hours.

[0098] A solution is prepared, consisting of 30.34 g of demineralized water and 8.0 g of copper nitrate, $\text{Cu}(\text{NO}_3)_2 \cdot 2.5\text{H}_2\text{O}$ (MW = 232.59, 34.4 mmoles). The solid is impregnated at room temperature under vacuum. It is left to slowly rotate, under vacuum for 3 h. It is then dried in an oven for 2 h at 120°C. It is calcined at 316°C/4h with an increase rate of 1°C min⁻¹.

[0099] A second impregnation, as the previous one but using a solution consisting of 4.06 g of Copper nitrate $\text{Cu}(\text{NO}_3)_2 \cdot 2.5\text{H}_2\text{O}$ (MW = 232.59, 17.5 mmoles) and 15.42 g of demineralized water, is effected on 18.8 g of product, calcined at 316°C.

[0100] The solid is impregnated at room temperature under vacuum. It is left to slowly rotate, under vacuum for 3 h. It is dried in an oven for 2 h at 120°C and is calcined at 482°C/8h with an increase rate of 1°C min⁻¹.

[0101] 19.4 g of calcined product are recovered (10.29% Cu).

EXAMPLE 5 - Catalytic test

[0102] A catalytic test is carried out, using an experimental apparatus as described below.

[0103] The experimental apparatus consists of tanks for the reagents benzene and acetone, feeding pumps of the reagents benzene and acetone, a mass meter for the flow-rate control of hydrogen coming from a cylinder, a static mixer of the reagents before their inlet into the reaction, a preheating unit of the reagents, a steel reactor situated inside an electric heating oven equipped with temperature regulation inside the oven and inside the reactor, a pressure regulation system inside the reactor by means of a pneumatic valve, a cooler of the reaction effluent and a collection system of the liquid and gaseous products.

[0104] The reactor, situated inside the heating oven, consists of a steel cylindrical tube, with a mechanical sealing system and internal diameter equal to about 2 cm.

[0105] Along the major axis of the reactor there is a thermometer trap having a diameter equal to 1 mm, containing a thermocouple which can run freely along the major axis of the reactor and consequently along the major axis of the catalytic bed.

[0106] The catalytic system prepared as described in Example 1 is charged into the reactor, in a quantity equal to 14.5 g, with a size ranging from 1 to 2 mm, for a total height of the catalytic bed equal to 8.5 cm.

[0107] A quantity of inert quartz material is charged over and under the catalytic bed for a height equal to 2 cm over and 2 cm under the catalytic bed.

[0108] The catalyst is then subjected to drying in a stream of nitrogen at a temperature inside the reactor equal to 160°C for about 1 hour, and a stream of 5.2 ml/min of low pressure hydrogen is subsequently fed for 60 minutes, followed by a stream of 15.8 ml/min at 180°C for 120 minutes and finally a stream of 23.6 ml/min at 200°C for 180 minutes, after which the hydrogen feeding is interrupted and the temperature of the reactor is brought back to a value equal to 150°C, the experimental apparatus being continuously maintained in a stream of nitrogen.

[0109] Once a constant temperature of 150°C has been reached, the stream of nitrogen is then interrupted and the feeding of benzene is initiated with a flow-rate equal to 0.245 ml/min.

[0110] The system is maintained under these conditions for 60 minutes, after which the feeding of hydrogen is reactivated at a flow-rate equal to 27.3 ml/min and after a few minutes the feeding of acetone is initiated with a flow-rate equal to 0.012 ml/min.

[0111] Approximately 3 hours after the feeding of acetone, samples of reaction effluent are removed, both for the liquid and gaseous part, which are subsequently analyzed by gaschromatography.

[0112] Table 1 summarizes the operating conditions, together with the results obtained. In this table:

- WHSV expresses the ratio between the sum of the hourly flow-rates of benzene and acetone (excluding hydrogen) and the quantity of catalytic system;

- the selectivity [aryls]/[acetone] expresses the fraction of acetone converted to cumene + polyisopropylbenzenes (products useful for the production of cumene in transalkylation) with respect to the total quantity of converted acetone;
- the selectivity [cumene]/[acetone] expresses the fraction of acetone converted to cumene with respect to the total quantity of converted acetone;
- the selectivity [aryls]/[benzene] expresses the fraction of benzene converted to cumene + polyisopropylbenzenes with respect to the total quantity of converted benzene.

EXAMPLE 6 - Catalytic test

[0113] A catalytic test is effected, using the same experimental apparatus and the same experimental conditions as Example 5, but charging the catalytic system prepared according to what is described in Example 2, in a quantity equal to 9 g, for a total height of the catalytic bed equal to 8.3 cm. The benzene is fed with a flow-rate equal to 0.184 ml/min.

[0114] Approximately 3 hours after the start of the acetone feeding, samples of reaction effluent are removed, both for the liquid and gaseous part, which are subsequently analyzed by gaschromatography.

[0115] Table 1 summarizes the operating conditions, together with the results obtained.

EXAMPLE 7 - Catalytic test

[0116] A catalytic test is effected, using the same experimental apparatus and the same experimental conditions as Example 5, but charging the catalytic system prepared according to what is described in Example 3, in a quantity equal to 11 g, for a total height of the catalytic bed equal to 8.5 cm. The benzene is fed with a flow-rate equal to 0.251 ml/min.

[0117] Approximately 3 hours after the start of the acetone feeding, samples of reaction effluent are removed, both for the liquid and gaseous part, which are subsequently analyzed by gaschromatography.

[0118] Table 1 summarizes the operating conditions, together with the results obtained.

EXAMPLE 8 - Catalytic test

[0119] A catalytic test is effected, using the same experimental apparatus and the same experimental conditions as Example 5, but charging the catalytic system prepared according to what is described in Example 3, in a quantity equal to 11 g, for a total height of the catalytic bed equal to 8.5 cm. The acetone is fed with a flow-rate equal to 0.009 ml/min, the benzene with a flow-rate equal to 0.072 ml/min.

[0120] Approximately 3 hours after the start of the acetone feeding, samples of reaction effluent are removed, both for the liquid and gaseous part, which are subsequently analyzed by gaschromatography.

[0121] Table 1 summarizes the operating conditions, together with the results obtained.

EXAMPLE 9 - Catalytic test

[0122] A catalytic test is effected, using the same experimental apparatus and the same experimental conditions as Example 5, but charging the catalytic system prepared according to what is described in Example 4, in a quantity equal to 5 g, for a total height of the catalytic bed equal to 5 cm.

[0123] The catalyst is then subjected to drying in a stream of nitrogen and the temperature inside the reactor is brought from 120°C to 190°C in about 2 hours. Once a constant temperature of 190°C has been reached, the stream of nitrogen is then interrupted and the feeding of benzene is initiated with a flow-rate equal to 0.254 ml/min. The system is maintained under these conditions for 60 minutes, after which the feeding of hydrogen is reactivated at a flow-rate equal to 27.3 ml/min and after a few minutes the feeding of acetone is initiated with a flow-rate equal to 0.036 ml/min.

[0124] Approximately 3 hours after the feeding of acetone, samples of reaction effluent are removed, both for the liquid and gaseous part, which are subsequently analyzed by gaschromatography. Table 1 summarizes the operating conditions, together with the results obtained.

TABLE 1

Example Nr.	5	6	7	8	9
Quantity of catalytic system (g)	14.5	9	11	11	5
Reaction temperature (°C)	150	150	150	150	190
Reaction pressure (kpa)	100	100	100	100	100
Total WHSV (h ⁻¹)	0.9	1.1	1.2	0.4	3.0

(continued)

Example Nr.	5	6	7	8	9
[Benz.]/[Acetone] molar ratio	16.8	12.6	17.2	6.3	5.8
[H ₂]/[Acetone] molar ratio	7.4	7.4	7.4	9.77	2.5
Conversion of acetone %	100.0	100	100	100	98.5
[aryls]/[acetone] selectivity %	96.6	88.3	94.6	98.0	89.1
[cumene]/[acetone] selectivity %	81.9	70.2	79.8	79.7	81.0
[aryls]/[benzene] selectivity %	99.4	96.7	99.0	99.8	96.7

EXAMPLE 10 - Preparation of the catalytic system

[0125] A material is prepared, consisting of a catalyst based on copper aluminate (produced by SüdChemie under the trade-name of T4489, indicated hereunder as "material A2", having the following composition expressed as weight percentage of the elements: Cu 39.3%, Al 15.5%, Zn 6.0%, Mn 6.8%), zeolite beta and alumina p-bohemite. The zeolite beta and alumina are the same materials used in the previous examples.

[0126] 173 g of alumina, 192 g of zeolite beta, 124 g of T4489 are charged into a ploughshare mixer and mechanically mixed for about 60 minutes, after which 400 cc of an aqueous solution of acetic acid at 2% w/w are added to the mixture of powders thus obtained, without interrupting the mixing for a further 60 minutes approximately.

[0127] The intermediate thus obtained is then subjected to extrusion using a HUTT type gear press extruder and the product obtained is then subjected to aging for a period of not less than 48 hours.

[0128] After the aging period, the product, in the form of pellets, is subsequently subjected to calcination treatment in air at a temperature of about 550°C for 5 hours, obtaining a material called "material B3", containing about 30% of copper aluminate T4489, about 40% of zeolite beta and about 30% of an alumina binding carrier. The catalytic system is prepared so that it consists of two distinct zones, the first containing 3 g of the material A2 and the second 7 g of the material B3 prepared as described. The total quantity of catalytic system is therefore equal to 10 g.

EXAMPLE 11 - Preparation of the catalytic system

[0129] A material consisting of copper aluminate T4489 and zeolite beta, already used in the previous example, is prepared.

[0130] 298 g of zeolite beta, 255 g of T4489 are charged into a ploughshare mixer and mechanically mixed for about 60 minutes, after which 365 cc of an aqueous solution of acetic acid at 5% w/w are added to the mixture of powders thus obtained, without interrupting the mixing for a further 60 minutes approximately.

[0131] The intermediate thus obtained is then subjected to extrusion using a HUTT type gear press extruder and the product obtained is then subjected to aging for a period of not less than 48 hours.

[0132] After the aging period, the product, in the form of pellets, is subsequently subjected to calcination treatment in air at a temperature of about 550°C for 5 hours, obtaining a material called "material B4", containing about 50% of copper aluminate T4489 and about 50% of zeolite beta. The catalytic system is prepared so that it consists of two distinct zones, the first containing 3 g of the material A2 and the second 7 g of the material B4 prepared as described. The total quantity of catalytic system is therefore equal to 10 g.

EXAMPLE 12 - Catalytic test

[0133] A catalytic test is carried out, using the same experimental apparatus described in Example 5.

[0134] The catalytic system prepared as described in Example 10 is charged into the reactor in a quantity equal to 10 g, with a size ranging from 1 to 2 mm, for a total height of the catalytic bed equal to 8.5 cm.

[0135] A quantity of inert quartz material is charged over and under the catalytic bed for a height equal to 2 cm over and 2 cm under the catalytic bed.

[0136] The catalyst is then subjected to drying in a stream of nitrogen for 1 hour during which the internal temperature is brought to 210°C. Once this temperature has been reached, a stream of 1.3 ml/min of benzene is fed for about 30 minutes. The temperature is then lowered to 194°C, the pressure is brought to 2100 Kpa and the nitrogen flow is interrupted. A stream of 367 ml/min of hydrogen is subsequently fed for about 30 minutes. Finally, the benzene feeding is interrupted and a flow of 0.77 g/min of a benzene and acetone solution [C6]/[C3] = 6.3, is fed.

[0137] About 3 hours after the start of the acetone feeding, samples of effluent are taken from the reaction, both for

the liquid and gaseous part, which are subsequently analyzed by means of gaschromatography. Approximately 24 hours after the start of the feeding, the sampling operation is repeated with the same analysis procedure.

[0138] Table 2 summarizes the operating conditions, together with the results obtained.

EXAMPLE 13 - Catalytic test

[0139] A catalytic test is carried out, using the same experimental apparatus described in Example 5 and under the same experimental conditions as Example 12. The catalytic system prepared as described in Example 11 is charged into the reactor in a quantity equal to 10 g, with a size ranging from 1 to 2 mm, for a total height of the catalytic bed equal to 8.1 cm.

[0140] About 3 hours after the start of the acetone feeding, samples of effluent are taken from the reaction, both for the liquid and gaseous part, which are subsequently analyzed by means of gaschromatography. Approximately 24 hours after the start of the feeding, the sampling operation is repeated with the same analysis procedure.

[0141] Table 2 summarizes the operating conditions, together with the results obtained.

EXAMPLE 14 - Catalytic test

[0142] A catalytic test is carried out, using the same experimental apparatus described in Example 5 and under the same experimental conditions as Example 12, but charging the catalytic system prepared as described in Example 3 in a quantity equal to 11 g for a total height of the catalytic bed equal to 8.3 cm. A flow of 0.76 g/min of a solution of benzene and acetone $[C_6]/[C_3] = 6.3$, is fed.

[0143] About 3 hours after the start of the acetone feeding, samples of effluent are taken from the reaction, both for the liquid and gaseous part, which are subsequently analyzed by means of gaschromatography. Approximately 24, 173 and 384 hours after the start of the feeding, the sampling operation is repeated with the same analysis procedure.

[0144] Table 2 summarizes the operating conditions, together with the results obtained.

EXAMPLE 15 - Catalytic test

[0145] A catalytic test is carried out, using the same experimental apparatus described in Example 5, but charging the catalytic system prepared as described in Example 3 in a quantity equal to 11 g for a total height of the catalytic bed equal to 8.4 cm.

[0146] The catalyst is subjected to drying in a stream of nitrogen at a temperature inside the reactor equal to 160°C for about 1 hour, after which a flow of 5.2 ml/min of hydrogen is fed at low pressure for 60 minutes, followed by a flow of 15.8 ml/min at 180°C for 120 minutes and finally a flow of 23.6 ml/min at 200°C for 180 minutes. The hydrogen feeding is then interrupted and the temperature of the reactor is brought to a value equal to 170°C, the experimental apparatus being maintained in a stream of nitrogen.

[0147] Once the temperature has reached a constant value of 170°C, the nitrogen stream is interrupted and the feeding of benzene is initiated with a flow-rate equal to 0.36 ml/min. The pressure is brought to 850 Kpa.

[0148] The system is maintained under these conditions for 60 minutes, after which the feeding of hydrogen is restarted at a flow-rate equal to 338 ml/min; after a few minutes, the benzene feeding is interrupted and a flow of 0.76 g/min of a solution of benzene and acetone $[C_6]/[C_3] = 6.3$, is fed.

[0149] About 3 hours after the start of the acetone feeding, samples of effluent are taken from the reaction, both for the liquid and gaseous part, which are subsequently analyzed by means of gaschromatography.

[0150] Table 2 summarizes the operating conditions, together with the results obtained.

EXAMPLE 16 - Catalytic test

[0151] A catalytic test is carried out, using the same experimental apparatus described in Example 5, but charging the catalytic system prepared as described in Example 3 in a quantity equal to 11 g for a total height of the catalytic bed equal to 8.4 cm.

[0152] The catalyst is subjected to drying in a stream of nitrogen for 1 hour during which the temperature inside the reactor is brought to 230°C. Once this temperature has been reached, a flow of 1.3 ml/min of benzene is fed for about 30 minutes. The temperature is then lowered to 210°C, the pressure is brought to 2900 Kpa, the nitrogen stream is interrupted and a flow of 367 ml/min of hydrogen is fed for about 30 minutes. Finally, the benzene feeding is interrupted and a flow of 0.76 g/min of a solution of benzene and acetone $[C_6]/[C_3] = 6.3$, is fed.

[0153] About 20 hours after the start of the acetone feeding, samples of effluent are taken from the reaction, both for the liquid and gaseous part, which are subsequently analyzed by means of gaschromatography.

[0154] Table 2 summarizes the operating conditions, together with the results obtained.

Table 2

Example Nr.	12		13		14				15		16
	3	24	3	24	3	24	173	384	3	24	20
Time of stream (h)	10	10	10	10	11	11	11	11	11	11	11
Quantity of catalytic system (g)	194	194	194	194	194	194	194	194	174	174	210
Reaction temperature (°C)	2100	2100	2100	2100	2100	2100	2100	2100	850	850	2900
Reaction pressure (kpa)	4.6	4.6	4.6	4.6	4.1	4.1	4.1	4.1	4.1	4.1	4.1
Total WHSV (h ⁻¹)	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3
[Benz.]/[Acetone] molar ratio	11.9	11.9	11.9	11.9	11.9	11.9	11.9	11.9	10.8	10.8	11.9
[H ₂]/[Acetone] molar ratio	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0
Conversion of acetone (%)	97.8	97.8	97.6	97.8	97.7	97.8	97.7	97.0	97.1	96.2	96.9
[Aryls]/[Acetone] selectivity (%)	76.6	76.7	73.9	73.7	76.9	77.0	76.8	76.2	68.7	70.1	83.8
[Cumene]/[Acetone] selectivity (%)	98.8	99.0	98.7	98.8	99.2	99.3	99.3	99.1	99.6	99.3	99.2
[Aryls]/[Benzene] selectivity (%)											

Claims

1. A process for preparing isopropylbenzene which comprises reacting benzene with acetone and hydrogen, in a single step, in the presence of a catalytic composition comprising a solid acid material and copper.
2. The process according to claim 1, wherein the catalytic composition contains one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides.
3. The process according to claim 2, wherein the catalytic composition contains one or more elements selected from elements of groups IIIA, and VIB.
4. The process according to claim 1, wherein the copper is in the form of an oxide.
5. The process according to claim 2, wherein the element selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, is in the form of an oxide.
6. The process according to claims 5, wherein copper and the elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, are contained in the catalytic composition in the form of mixed oxide.
7. The process according to claim 3, wherein the element selected from elements of groups IIIA and VIB is in the form of an oxide.
8. The process according to claim 7, wherein copper and the elements of groups IIIA and VIB are contained in the catalytic composition in the form of a mixed oxide.
9. The process according to claim 3 or 7, wherein the element of group VIB or IIIA is selected from Cr and Al.
10. The process according to claim 9, wherein the catalytic composition contains Cr and Cu in the form of copper chromite.
11. The process according to one or more of the previous claims, wherein the solid acid material is selected from one or more zeolitic materials in acid or prevalently acid form.
12. The process according to claim 11, wherein the zeolitic material is selected from zeolite beta, zeolite Y, ZSM-12 and mordenite.
13. The process according to claim 12, wherein the zeolite is zeolite beta.
14. The process according to claim 13, wherein the catalytic composition comprises copper chromite and zeolite beta.
15. The process according to claim 1, wherein the copper is contained in the catalytic composition in a weight ratio of the metal with respect to the solid acid material ranging from 0.001 to 10.
16. The process according to claim 15, wherein the copper is contained in the catalytic composition in a weight ratio of the metal with respect to the solid acid material ranging from 0.01 to 2.
17. The process according to claim 2, wherein the element selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, is contained in the catalytic composition in a weight ratio of the metal with respect to the solid acid material ranging from 0.001 to 10.
18. The process according to claim 17, wherein the element selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, is contained in the catalytic composition in a weight ratio of the metal with respect to the solid acid material ranging from 0.01 to 2.
19. The process according to claim 10, wherein the catalytic composition comprises promoters such as Ba and Mn.

20. The process according to claim 19, wherein Ba or Mn are contained in the catalytic composition in a weight percentage ranging from 0.1 to 5% with respect to the weight of the composition.
21. The process according to one or more of the previous claims, wherein the catalytic composition comprises an inorganic binder.
22. The process according to claim 21, wherein the binder is selected from silicon oxide and aluminum oxide.
23. The process according to claim 22, wherein the binder is γ -alumina.
24. The process according to any of the previous claims, wherein the catalytic composition consists of one or more distinct zone each of which contain a metallic component, containing copper and optionally one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, or a solid acid component, or both components.
25. The process according to claim 24, wherein the zone containing the metallic component is made up of copper oxide, or copper chromite, or is a mixed oxide of copper and one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides.
26. The process according to claim 24, wherein the zone containing the solid acid component is of a zeolitic nature.
27. The process according to claim 26, wherein the zeolite is selected from zeolite beta, zeolite Y, ZSM-12 and mordenite.
28. The process according to claim 27, wherein the zeolite is zeolite beta.
29. The process according to claim 24, wherein the zone containing both components is prepared by impregnation of the acid solid with a solution of a copper salt and optionally a salt of one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, drying and calcination.
30. The process according to claim 29, wherein the composition containing both components is used in a mixture with a binding agent according to claims 21, 22 or 23.
31. The process according to claim 24, wherein the zone containing both components is prepared by impregnation of the mixture of the solid acid material and a binding agent according to claims 21, 22 or 23, with a solution of a copper salt and optionally a salt of one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, drying and calcination.
32. The process according to claim 24, wherein the zone containing both components is prepared by means of an ion exchange process in which the solid acid material is put in an aqueous solution containing a copper salt and optionally a salt of one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, the mixture is left under stirring for several hours, the solid in suspension is recovered by filtration, washed with demineralized water and dried, obtaining a solid acid material in exchanged form.
33. The process according to claim 30, wherein the material containing both components is used in a mixture with binding agents according to claims 21, 22 or 23.
34. The process according to claim 24, wherein the zone containing both components is prepared by means of an ion exchange process in which a composition of the solid acid material and binding agents according to claims 21, 22 or 23 is put in an aqueous solution containing a copper salt and optionally a salt of one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, group VIII limited to Fe, Ru and Os, and of the series of lanthanides, the mixture is left under stirring for several hours, the solid in suspension is recovered by filtration, washed with demineralized water and dried.
35. The process according to claim 24, wherein the zone containing both components is prepared by means of the extrusion, drying and calcination of a mechanical mixture of the two components paste-mixed with a peptizing agent.

36. The process according to claim 24, wherein the zone containing both components is prepared by means of the extrusion, drying and calcination of a mechanical mixture of the solid acid component, the metallic component and binding agents according to claims 21, 22 or 23.
- 5 37. The process according to claim 24, wherein the catalytic composition consists of a single zone containing both the metallic and solid acid components.
38. The process according to claim 24, wherein the catalytic composition consists of two or more distinct zones each containing both the metallic and solid acid components, wherein the composition of the single zones is different.
- 10 39. The process according to claim 24 or 25, wherein the catalytic composition consists of two or more distinct zones, one of which containing the metallic component, whereas the others contain one or more different combinations of the two metallic and solid acid components.
- 15 40. The process according to claim 24 or 25, wherein the catalytic composition consists of two or more distinct zones, one of which containing the metallic component, whereas the subsequent zones contain the solid acid component.
41. The process according to claim 24 or 25, wherein the catalytic composition consists of a single zone containing the two components of the catalytic composition, mechanically mixed with each other.
- 20 42. The process according to claim 24 or 25, wherein the catalytic composition consists of two or more distinct zones, one of which containing the metallic component, whereas the others contain different pairs of the two metallic and solid acid components, mechanically mixed.
- 25 43. The process according to claim 39, 40 or 42, wherein the zone containing the metallic component is that which first comes into contact with the stream of reagents consisting of the aromatic compound, ketone and hydrogen.
44. The process according to claim 24, wherein the catalytic composition contains the metallic component with an activity gradient which decreases in one direction and the solid acid component with an activity gradient which decreases in the opposite direction.
- 30 45. The process according to claim 44, wherein the activity gradient of the metallic component decreases along the feeding direction and flow of the reagents, whereas the activity gradient of the solid acid component decreases along the opposite direction and flow.
- 35 46. The process according to claim 1 or 2, carried out at a temperature ranging from 50 to 350°C and at a pressure equal to or higher than the atmospheric pressure.
- 40 47. The process according to claim 46, carried out at a temperature ranging from 100 to 250°C and at a pressure ranging from 1 to 50 bars.
48. The process according to claim 1 or 2, wherein a molar ratio not lower than 1 is used in the feeding between the aromatic hydrocarbon and ketone.
- 45 49. The process according to claim 48, wherein the molar ratio is higher than 2.
50. The process according to claim 1 or 2, wherein a molar ratio not lower than 1 is used between hydrogen and ketone in the feeding.
- 50 51. The process according to claim 50, wherein a molar ratio higher than 2 is used between hydrogen and ketone in the feeding.
- 55 52. The process according to any of the previous claims, wherein the aromatic compound is benzene, the ketone is acetone and the product resulting from the alkylation process, which contains isopropylbenzene, non-converted benzene, non-converted hydrogen, water and polyisopropylbenzenes, is fractionated in a separation section to obtain a first fraction mainly containing hydrogen, a second fraction mainly containing water, a third fraction mainly containing benzene, a fourth fraction mainly containing isopropylbenzene and a fifth fraction mainly containing polyisopropylbenzenes, wherein the first fraction is re-used in the reaction step with acetone and benzene, the

second fraction is removed from the process, the third fraction is partly re-used in the reaction step with acetone and hydrogen and partly in a subsequent transalkylation step, where it is reacted with the fifth fraction to product isopropylbenzene.

53. The process according to claim 52, wherein the transalkylation reaction product is fractionated into benzene, which is re-fed to the alkylation process, cumene and polyisopropylbenzenes, wherein the polyisopropylbenzenes are re-fed to the transalkylation step.

54. A process for the production of phenol which comprises the following steps:

(a) reacting benzene, acetone and hydrogen in the presence of a catalytic system comprising a solid acid material, copper, and optionally one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides,

(b) oxidization of the cumene to cumene hydroperoxide,

(c) rearrangement of the cumene hydroperoxide to phenol and acetone,

characterized in that step (a) is effected according to one or more of claims 1 to 53.

55. The process according to claim 54, wherein the acetone which is formed in step (c) is recycled to the synthesis step (a) of cumene.

56. A catalytic composition containing a zeolite in acid form, Cu and one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides, with the proviso that when zeolite is zeolite Y and the catalytic composition contains one or more elements of Group VIB, said element is selected between Mo and W.

57. The catalytic composition according to claim 56 containing a zeolite in acid form selected among zeolite beta, zeolite ZSM-12 and mordenite, Cu and one or more elements selected from elements of Groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of Group VIII limited to Fe, Ru and Os, and of the series of lanthanides.

58. The catalytic composition according to claim 56 containing zeolite Y in acid form, Cu and one or more elements selected from elements of Groups IIIA, IVA, IIIB, IVB, VB, VIIB, of Group VIII limited to Fe, Ru and Os, and of the series of lanthanides.

59. The catalytic composition according to claim 56, containing a binder.

60. The catalytic composition according to claim 56 or 59, wherein the zeolite is zeolite beta.

61. The catalytic composition according to claim 60, containing zeolite beta, copper, an element selected from Cr and Al, and optionally a binder.

62. The catalytic composition according to claim 61, wherein copper and chromium are in the form of copper chromite.

63. A process for preparing the catalytic composition according to claim 56, comprising the impregnation of the zeolite with an aqueous solution containing a copper salt and a salt of one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides, drying and calcination.

64. The process according to claim 63, wherein the resulting material is bound with a binder according to claims 21, 22 or 23.

65. A process for preparing the catalytic composition according to claim 56, comprising the impregnation of the zeolite and binding agents according to claims 21, 22 or 23 with an aqueous solution containing a copper salt and a salt of one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides, drying and calcination.

66. A process for preparing the catalytic composition according to claim 56, comprising the ionic exchange of the zeolite with an aqueous solution containing a copper salt and a salt of one or more elements selected from elements of

groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides, drying and calcination.

67. The process for preparing the catalytic composition according to claim 66, wherein the material obtained is bound with a binder.

68. A process for preparing the catalytic composition according to claim 56, comprising the ionic exchange of a composition of zeolite and suitable binding agents with an aqueous solution containing a copper salt and a salt of one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides, drying and calcination.

69. A process for preparing the catalytic composition according to claim 56, comprising the extrusion, drying and calcination of a mechanical mixture, paste-mixed with a peptizing agent, of zeolite, Cu and one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides, in the form of oxides.

70. A process for preparing the catalytic composition according to claim 56, comprising the extrusion, drying and calcination of a mechanical mixture, paste-mixed with a peptizing agent, of zeolite, Cu and one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides, in the form of oxides, and binding agents according to claims 21, 22 or 23.

71. A process for preparing the catalytic composition according to claim 56, which comprises assembling various distinct zone each containing the metallic component containing copper and one or more elements selected from elements of groups IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, of group VIII limited to Fe, Ru and Os, and of the series of lanthanides, or the zeolitic component, or both components.

Patentansprüche

1. Verfahren zum Herstellen von Isopropylbenzol, umfassend Reagieren von Benzol mit Aceton und Wasserstoff, in einem einzelnen Schritt, in der Gegenwart einer katalytischen Zusammensetzung, umfassend ein Feststoffsäurematerial und Kupfer.

2. Verfahren nach Anspruch 1, wobei die katalytische Zusammensetzung ein oder mehrere Element(e), ausgewählt aus den Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, enthält.

3. Verfahren nach Anspruch 2, wobei die katalytische Zusammensetzung ein oder mehrere Element(e), ausgewählt aus den Elementen von Gruppen IIIA und VIB, enthält.

4. Verfahren nach Anspruch 1, wobei das Kupfer in der Form eines Oxids vorliegt.

5. Verfahren nach Anspruch 2, wobei das Element ausgewählt aus Elementen von den Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und aus der Lanthanoid-Reihe, in der Form eines Oxids vorliegt.

6. Verfahren nach Anspruch 5, wobei Kupfer und die Elemente von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe in der katalytischen Zusammensetzung in der Form von gemischten Oxiden, enthalten sind.

7. Verfahren nach Anspruch 3, wobei das Element, ausgewählt aus Elementen von Gruppen IIIA und IVB, in der Form von einem Oxid vorliegt.

8. Verfahren nach Anspruch 7, wobei Kupfer und die Elemente von Gruppen IIIA und VIB in der katalytischen Zusammensetzung in der Form eines gemischten Oxids enthalten sind.

9. Verfahren nach Anspruch 3 oder 7, wobei das Element von Gruppe VIB oder IIIA aus Cr und Al ausgewählt ist.

10. Verfahren nach Anspruch 9, wobei die katalytische Zusammensetzung Cr und Cu in der Form von Kupfer-Chromit enthält.
- 5 11. Verfahren nach einem oder mehreren der vorhergehenden Ansprüche, wobei das Feststoffsäurematerial ausgewählt ist aus einem oder mehreren zeolithischen Materialien in saurer oder vorherrschend saurer Form.
12. Verfahren nach Anspruch 11, wobei das zeolithische Material ausgewählt ist aus Zeolith beta, Zeolith Y, ZSM-12 und Mordenit.
- 10 13. Verfahren nach Anspruch 12, wobei das Zeolith Zeolith beta ist.
14. Verfahren nach Anspruch 13, wobei die katalytische Zusammensetzung Kupfer-Chromit und Zeolith beta umfasst.
- 15 15. Verfahren nach Anspruch 1, wobei das Kupfer in der katalytischen Zusammensetzung in einem Gewichtsverhältnis des Metalls, bezogen auf das Feststoffsäurematerial, im Bereich von 0,001 bis 10 enthalten ist.
16. Verfahren nach Anspruch 15, wobei das Kupfer in der katalytischen Zusammensetzung in einem Gewichtsverhältnis des Metalls, bezogen auf das Feststoffsäurematerial, im Bereich von 0,01 bis 2 enthalten ist.
- 20 17. Verfahren nach Anspruch 2, wobei das Element ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und aus der Lanthanoid-Reihe in der katalytischen Zusammensetzung in einem Gewichtsverhältnis des Metalls, bezogen auf das Feststoffsäurematerial, im Bereich von 0,001 bis 10, enthalten ist.
- 25 18. Verfahren nach Anspruch 17, wobei das Element ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und aus der Lanthanoid-Reihe in der katalytischen Zusammensetzung in einem Gewichtsverhältnis des Metalls, bezogen auf das Feststoffsäurematerial, im Bereich von 0,01 bis 2, enthalten ist.
- 30 19. Verfahren nach Anspruch 10, wobei die katalytische Zusammensetzung Promotoren wie Ba und Mn umfasst.
20. Verfahren nach Anspruch 19, wobei Ba oder Mn in der katalytischen Zusammensetzung enthalten sind im Bereich von 0,1 bis 5 Gewichtsprozent, bezogen auf das Gewicht der Zusammensetzung.
- 35 21. Verfahren nach einem oder mehreren der vorhergehenden Ansprüche, wobei die katalytische Zusammensetzung ein anorganisches Bindemittel umfasst.
22. Verfahren nach Anspruch 21, wobei das Bindemittel ausgewählt ist aus Siliziumoxid und Aluminiumoxid.
- 40 23. Verfahren nach Anspruch 22, wobei das Bindemittel γ -Aluminiumoxid ist.
24. Verfahren nach einem beliebigen der vorhergehenden Ansprüche, wobei die katalytische Zusammensetzung aus einer oder mehreren getrennten Zonen besteht, von denen jede eine metallische Komponente enthält, enthaltend Kupfer und gegebenenfalls eines oder mehrere Elemente, ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, oder eine Feststoff-
45 säurekomponente, oder beide Komponenten.
25. Verfahren nach Anspruch 24, wobei die Zone, die eine metallische Komponente enthält, zusammengesetzt ist aus Kupferoxid oder Kupfer-Chromit, oder ein gemischtes Oxid von Kupfer und einem oder mehreren Elementen, ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und
50 Os, und der Lanthanoid-Reihe, ist.
26. Verfahren nach Anspruch 24, wobei die Zone, die die Feststoffsäurekomponente enthält, von zeolithischer Natur ist.
- 55 27. Verfahren nach Anspruch 26, wobei das Zeolith ausgewählt ist aus Zeolith beta, Zeolith Y, ZSM-12 und Mordenit.
28. Verfahren nach Anspruch 27, wobei das Zeolith Zeolith beta ist.

29. Verfahren nach Anspruch 24, wobei die Zone, die beide Komponenten enthält, hergestellt wird durch Imprägnieren der Feststoffsäure mit einer Lösung eines Kupfersalzes und gegebenenfalls einem Salz von einem oder mehreren Elementen, ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, Trocknen und Kalzinieren.
30. Verfahren nach Anspruch 29, wobei die Zusammensetzung, die beide Komponenten enthält, in einer Mischung mit einem Bindemittel gemäß den Ansprüchen 21, 22 oder 23 verwendet wird.
31. Verfahren nach Anspruch 24, wobei die Zone, die beide Komponenten enthält, hergestellt wird durch Imprägnieren der Mischung des Feststoffsäurematerials und eines Bindemittels gemäß den Ansprüchen 21, 22 oder 23, mit einer Lösung eines Kupfersalzes und gegebenenfalls einem Salz von einem oder mehreren Elementen, ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, Trocknen und Kalzinieren.
32. Verfahren nach Anspruch 24, wobei die Zone, die beide Komponenten enthält, hergestellt wird mittels eines Ionenaustauschverfahrens, in dem das Feststoffsäurematerial in eine wässrige Lösung, enthaltend ein Kupfersalz und gegebenenfalls ein Salz von einem oder mehreren Elementen, ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, eingeführt wird, die Mischung für mehrere Stunden gerührt wird, der Feststoff in Suspension durch Filtrieren rückgewonnen wird, mit entmineralisiertem Wasser gewaschen wird und getrocknet wird, wodurch ein Feststoffsäurematerial in ausgetauschter Form erhalten wird.
33. Verfahren nach Anspruch 30, wobei das Material, das beide Komponenten enthält, in einer Mischung mit Bindemitteln gemäß den Ansprüchen 21, 22 oder 23 verwendet wird.
34. Verfahren nach Anspruch 24, wobei die Zone, die beide Komponenten enthält, hergestellt wird mittels eines Ionenaustauschverfahrens, in dem eine Zusammensetzung des Feststoffsäurematerials und Bindemitteln gemäß den Ansprüchen 21, 22 oder 23 in eine wässrige Lösung, enthaltend ein Kupfersalz und gegebenenfalls ein Salz von einem oder mehreren Elementen, ausgewählt aus den Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, eingeführt wird, die Mischung für mehrere Stunden rühren gelassen wird, der Feststoff in Suspension durch Filtrieren rückgewonnen wird, mit entmineralisiertem Wasser gewaschen wird und getrocknet wird.
35. Verfahren nach Anspruch 24, wobei die Zone, die beide Komponenten enthält, hergestellt wird mittels der Extrusion, dem Trocknen und dem Kalzinieren einer mechanischen Mischung der mit einem Mastifizierungsmittel Pasten-gemischten zwei Komponenten.
36. Verfahren nach Anspruch 24, wobei die Zone, die beide Komponenten enthält, hergestellt wird mittels der Extrusion, dem Trocknen und dem Kalzinieren einer mechanischen Mischung der Feststoffsäurekomponente, der metallischen Komponente und Bindemitteln gemäß den Ansprüchen 21, 22 oder 23.
37. Verfahren nach Anspruch 24, wobei die katalytische Zusammensetzung aus einer einzelnen Zone besteht, enthaltend sowohl die metallischen als auch die Feststoffsäurekomponenten.
38. Verfahren nach Anspruch 24, wobei die katalytische Zusammensetzung aus zwei oder mehr getrennten Zonen besteht, jede enthaltend sowohl die metallischen als auch die Feststoffsäurekomponenten, wobei die Zusammensetzung der einzelnen Zonen unterschiedlich ist.
39. Verfahren nach Anspruch 24 oder 25, wobei die katalytische Zusammensetzung aus zwei oder mehr getrennten Zonen besteht, von denen eine die metallische Komponente enthält, während die anderen eine oder mehrere unterschiedliche Kombinationen der zwei metallischen und Feststoffsäurekomponenten enthalten.
40. Verfahren nach Anspruch 24 oder 25, wobei die katalytische Zusammensetzung aus zwei oder mehr getrennten Zonen besteht, von denen eine die metallische Komponente enthält, während die folgenden Zonen die Feststoffsäurekomponente enthalten.
41. Verfahren nach Anspruch 24 oder 25, wobei die katalytische Zusammensetzung aus einer einzelnen Zone besteht, enthaltend die zwei Komponenten der katalytischen Zusammensetzung, die miteinander mechanisch gemischt sind.

42. Verfahren nach Anspruch 24 oder 25, wobei die katalytische Zusammensetzung aus zwei oder mehr getrennten Zonen besteht, von denen eine die metallische Komponente enthält, während die andere unterschiedliche Paare von zwei metallischen und Feststoffsäurekomponenten enthält, die mechanisch gemischt sind.
- 5 43. Verfahren nach Anspruch 39, 40 oder 42, wobei die Zone, enthaltend die metallische Komponente, diejenige ist, die als erstes mit dem Reagenzienstrom, bestehend aus der aromatischen Verbindung, dem Keton und dem Wasserstoff, in Kontakt kommt.
- 10 44. Verfahren nach Anspruch 24, wobei die katalytische Zusammensetzung die metallische Komponente mit einem Aktivitätsgradienten, der in eine Richtung abnimmt und die Feststoffsäurekomponente mit einem Aktivitätsgradienten, der in die entgegengesetzte Richtung abnimmt, enthält.
- 15 45. Verfahren nach Anspruch 44, wobei der Aktivitätsgradient der metallischen Zusammensetzung entlang der Zufuhr-richtung und dem Reagenzienfluss abnimmt, während der Aktivitätsgradient der Feststoffsäurekomponente entlang der entgegengesetzten Richtung und dem Fluss abnimmt.
46. Verfahren nach Anspruch 1 oder 2, durchgeführt bei einer Temperatur im Bereich von 50 bis 350 °C und bei einem Druck gleich oder höher als der Atmosphärendruck.
- 20 47. Verfahren nach Anspruch 46, durchgeführt bei einer Temperatur im Bereich von 100 bis 250 °C und bei einem Druck im Bereich von 1 bis 50 bar.
48. Verfahren nach Anspruch 1 oder 2, wobei ein molares Verhältnis nicht niedriger als 1 in der Zufuhr zwischen dem aromatischen Kohlenwasserstoff und dem Keton verwendet wird.
- 25 49. Verfahren nach Anspruch 48, wobei ein molares Verhältnis mehr als 2 ist.
50. Verfahren nach Anspruch 1 oder 2, wobei ein molares Verhältnis nicht niedriger als 1 zwischen Wasserstoff und Keton in der Zufuhr verwendet wird.
- 30 51. Verfahren nach Anspruch 50, wobei ein molares Verhältnis von mehr als 2 zwischen Wasserstoff und Keton in der Zufuhr verwendet wird.
- 35 52. Verfahren nach einem beliebigen der vorhergehenden Ansprüche, wobei die aromatische Verbindung Benzol ist, das Keton Aceton ist, und das Produkt resultierend aus dem Alkylierungsprozess, der Isopropylbenzol, nicht-konvertiertes Benzol, nicht-konvertierten Wasserstoff, Wasser und Polyisopropylbenzole enthält, in einem Trennungsabschnitt fraktioniert wird, um eine erste Fraktion hauptsächlich enthaltend Wasserstoff, eine zweite Fraktion hauptsächlich enthaltend Wasser, eine dritte Fraktion hauptsächlich enthaltend Benzol, eine vierte Fraktion hauptsächlich enthaltend Isopropylbenzol, und eine fünfte Fraktion hauptsächlich enthaltend Polyisopropylbenzole, zu erhalten,
- 40 wobei die erste Fraktion im Reaktionsschritt mit Aceton und Benzol wieder verwendet wird, die zweite Fraktion aus dem Verfahren entfernt wird, die dritte Fraktion teilweise in dem Reaktionsschritt mit Aceton und Wasserstoff wieder verwendet und teilweise in einem nachfolgenden Transalkylierungsschritt wieder verwendet wird, wo sie mit der fünften Fraktion reagiert wird, um Isopropylbenzol zu produzieren.
- 45 53. Verfahren nach Anspruch 52, wobei das Transalkylierungsreaktionsprodukt fraktioniert wird in Benzol, das wieder dem Alkylierungsverfahren zugeführt wird, Cumol und Polyisopropylbenzolen, wobei die Polyisopropylbenzole zu dem Transalkylierungsschritt wieder zugeführt werden.
- 50 54. Verfahren zum Herstellen von Phenol, umfassend die folgenden Schritte:
- (a) Reagieren von Benzol, Aceton und Wasserstoff in der Gegenwart eines katalytischen Systems, umfassend ein Feststoffsäurematerial, Kupfer und gegebenenfalls ein oder mehr Element(e), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe,
- 55 (b) Oxidieren des Cumols zu Cumolhydroperoxid,
- (c) Umlagern des Cumolhydroperoxids zu Phenol und Aceton,

dadurch gekennzeichnet, dass Schritt (a) gemäß einem oder mehreren der Ansprüche 1 bis 53 durchgeführt wird.

55. Verfahren nach Anspruch 54, wobei das Aceton, das in Schritt (c) gebildet wird, zu dem Syntheseschritt (a) von Cumol recycelt wird.
- 5 56. Katalytische Zusammensetzung, enthaltend ein Zeolith in saurer Form, Cu und ein oder mehrere Element(e), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, mit der Bedingung, dass wenn Zeolith Zeolith Y ist und die katalytische Zusammensetzung ein oder mehrere Element(e) von Gruppe VIB enthält, dieses Element zwischen Mo und W ausgewählt ist.
- 10 57. Katalytische Zusammensetzung nach Anspruch 56, enthaltend ein Zeolith in saurer Form, ausgewählt zwischen Zeolith beta, Zeolith ZSM-12 und Mordenit, Cu und einem oder mehreren Element(en), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe.
- 15 58. Katalytische Zusammensetzung nach Anspruch 56, enthaltend Zeolith Y in saurer Form, Cu und ein oder mehrere Element(e), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe.
59. Katalytische Zusammensetzung nach Anspruch 56, enthaltend ein Bindemittel.
- 20 60. Katalytische Zusammensetzung nach Anspruch 56 oder 59, wobei das Zeolith Zeolith beta ist.
61. Katalytische Zusammensetzung nach Anspruch 60, enthaltend Zeolith beta, Kupfer, ein Element ausgewählt aus Cr und Al, und gegebenenfalls ein Bindemittel.
- 25 62. Katalytische Zusammensetzung nach Anspruch 61, wobei Kupfer und Chrom in der Form von Kupfer-Chromit vorliegen.
- 30 63. Verfahren zum Herstellen der katalytischen Zusammensetzung nach Anspruch 56, umfassend das Imprägnieren des Zeoliths mit einer wässrigen Lösung, enthaltend ein Kupfersalz und ein Salz von einem oder mehreren Element(en), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, Trocknen und Kalzinieren.
- 35 64. Verfahren nach Anspruch 63, wobei das resultierende Material mit einem Bindemittel gemäß den Ansprüchen 21, 22 oder 23 gebunden wird.
- 40 65. Verfahren zum Herstellen der katalytischen Zusammensetzung nach Anspruch 56, umfassend das Imprägnieren des Zeoliths und Bindemitteln gemäß den Ansprüchen 21, 22 oder 23, mit einer wässrigen Lösung, enthaltend ein Kupfersalz und ein Salz von einem oder mehreren Element(en), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, Trocknen und Kalzinieren.
- 45 66. Verfahren zum Herstellen der katalytischen Zusammensetzung nach Anspruch 56, umfassend den Ionenaustausch des Zeoliths mit einer wässrigen Lösung, enthaltend ein Kupfersalz und ein Salz von einem oder mehreren Element(en), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, Trocknen und Kalzinieren.
- 50 67. Verfahren zum Herstellen der katalytischen Zusammensetzung nach Anspruch 66, wobei das erhaltene Material mit einem Bindemittel gebunden wird.
- 55 68. Verfahren zum Herstellen der katalytischen Zusammensetzung nach Anspruch 56, umfassend den Ionenaustausch einer Zusammensetzung von Zeolith und geeigneten Bindemitteln mit einer wässrigen Lösung, enthaltend ein Kupfersalz und ein Salz von einem oder mehreren Element(en), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, Trocknen und Kalzinieren.
69. Verfahren zum Herstellen der katalytischen Zusammensetzung nach Anspruch 56, umfassend das Extrudieren, Trocknen und Kalzinieren einer mechanischen Mischung, Pasten-gemischt mit einem Mastifizierungsmittel, von Zeolith, Cu und einem oder mehreren Element(en), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB,

VIIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, in der Form von Oxiden.

70. Verfahren zum Herstellen der katalytischen Zusammensetzung nach Anspruch 56, umfassend das Extrudieren, Trocknen und Kalzinieren einer mechanischen Mischung, Pasten-gemischt mit einem Mastifizierungsmittel, von Zeolith, Cu und einem oder mehreren Element(en), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, in der Form von Oxiden, und Bindemitteln gemäß den Ansprüchen 21, 22 oder 23.

71. Verfahren zum Herstellen der katalytischen Zusammensetzung nach Anspruch 56, umfassend Zusammensetzen unterschiedlicher getrennter Zonen, jede enthaltend die metallische Komponente, enthaltend Kupfer und eines oder mehrere Element(e), ausgewählt aus Elementen von Gruppen IIIA, IVA, IIIB, IVB, VB, VIB, VIIIB, Gruppe VIII, eingeschränkt auf Fe, Ru und Os, und der Lanthanoid-Reihe, oder die zeolithische Komponente, oder beide Komponenten.

Revendications

1. Procédé de préparation d'isopropylbenzène qui comprend la réaction du benzène avec de l'acétone et de l'hydrogène, dans une seule étape, en présence d'une composition catalytique comprenant un matériau acide solide et du cuivre.

2. Procédé selon la revendication 1, dans lequel la composition catalytique contient un ou plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides.

3. Procédé selon la revendication 2, dans lequel la composition catalytique contient un ou plusieurs éléments choisis parmi des éléments des groupes IIIA et VIB.

4. Procédé selon la revendication 1, dans lequel le cuivre est sous la forme d'un oxyde.

5. Procédé selon la revendication 2, dans lequel l'élément choisi parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, est sous la forme d'un oxyde.

6. Procédé selon la revendication 5, dans lequel le cuivre et les éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, sont contenus dans la composition catalytique sous la forme d'un oxyde mixte.

7. Procédé selon la revendication 3, dans lequel l'élément choisi parmi des éléments des groupes IIIA et VIB est sous la forme d'un oxyde.

8. Procédé selon la revendication 7, dans lequel le cuivre et les éléments des groupes IIIA et VIB sont contenus dans la composition catalytique sous la forme d'un oxyde mixte.

9. Procédé selon les revendications 3 ou 7, dans lequel l'élément du groupe VIB ou IIIA est choisi parmi Cr et Al.

10. Procédé selon la revendication 9, dans lequel la composition catalytique contient Cr et Cu sous la forme de chromite de cuivre.

11. Procédé selon une ou plusieurs des revendications précédentes, dans lequel le matériau acide solide est choisi parmi un ou plusieurs matériaux zéolithiques sous forme acide ou principalement acide.

12. Procédé selon la revendication 11, dans lequel le matériau zéolithique est choisi parmi la zéolite beta, la zéolite Y, ZSM-12 et la mordénite.

13. Procédé selon la revendication 12, dans lequel la zéolite est la zéolite beta.

14. Procédé selon la revendication 13, dans lequel la composition catalytique comprend du chromite de cuivre et de la zéolite beta.

15. Procédé selon la revendication 1, dans lequel le cuivre est contenu dans la composition catalytique dans un rapport pondéral du métal par rapport au matériau acide solide situé dans la plage de 0,001 à 10.
- 5 16. Procédé selon la revendication 15, dans lequel le cuivre est contenu dans la composition catalytique dans un rapport pondéral du métal par rapport au matériau acide solide situé dans la plage de 0,01 à 2.
- 10 17. Procédé selon la revendication 2, dans lequel l'élément choisi parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, est contenu dans la composition catalytique dans un rapport pondéral du métal par rapport au matériau acide solide situé dans la plage de 0,001 à 10.
- 15 18. Procédé selon la revendication 17, dans lequel l'élément choisi parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, est contenu dans la composition catalytique dans un rapport pondéral du métal par rapport au matériau acide solide situé dans la plage de 0,01 à 2.
- 20 19. Procédé selon la revendication 10, dans lequel la composition catalytique comprend des promoteurs tels que Ba et Mn.
- 25 20. Procédé selon la revendication 19, dans lequel Ba ou Mn sont contenus dans la composition catalytique dans un pourcentage pondéral situé dans la plage de 0,1 à 5 % par rapport au poids de la composition.
- 30 21. Procédé selon une ou plusieurs des revendications précédentes, dans lequel la composition catalytique comprend un liant inorganique.
22. Procédé selon la revendication 21, dans lequel le liant est choisi parmi l'oxyde de silicium et l'oxyde d'aluminium.
23. Procédé selon la revendication 22, dans lequel le liant est la γ -alumine.
- 35 24. Procédé selon l'une quelconque des revendications précédentes, dans lequel la composition catalytique est constituée d'une ou de plusieurs zones distinctes dont chacune contient un composant métallique, contenant du cuivre et éventuellement un ou plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, ou un composant acide solide, ou les deux composants.
- 40 25. Procédé selon la revendication 24, dans lequel la zone contenant le composant métallique est constitué d'oxyde de cuivre, ou de chromite de cuivre, ou est un oxyde mixte de cuivre et d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides.
- 45 26. Procédé selon la revendication 24, dans lequel la zone contenant le composant acide solide est d'une nature zéolitique.
27. Procédé selon la revendication 26, dans lequel la zéolite est choisie parmi la zéolite beta, la zéolite Y, ZSM-12 et la mordenite.
- 50 28. Procédé selon la revendication 27, dans lequel la zéolite est la zéolite beta.
29. Procédé selon la revendication 24, dans lequel la zone contenant les deux composants est préparée par imprégnation du solide acide avec une solution d'un sel de cuivre et éventuellement d'un sel d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, séchage et calcination.
- 55 30. Procédé selon la revendication 29, dans lequel la composition contenant les deux composants est utilisée en mélange avec un liant selon les revendications 21, 22 ou 23.
31. Procédé selon la revendication 24, dans lequel la zone contenant les deux composants est préparée par imprégnation du mélange du matériau acide solide et d'un liant selon les revendications 21, 22 ou 23, avec une solution d'un sel de cuivre et éventuellement d'un sel d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, séchage et calcination.

32. Procédé selon la revendication 24, dans lequel la zone contenant les deux composants est préparée au moyen d'un procédé d'échange d'ions dans lequel le matériau acide solide est introduit dans une solution aqueuse contenant un sel de cuivre et éventuellement un sel d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, le mélange est laissé sous agitation pendant plusieurs heures, le solide en suspension est récupéré par filtration, lavé avec de l'eau déminéralisée et séché, pour obtenir un matériau acide solide sous forme échangée.
33. Procédé selon la revendication 30, dans lequel le matériau contenant les deux composants est utilisé en mélange avec des liants selon les revendications 21, 22 ou 23.
34. Procédé selon la revendication 24, dans lequel la zone contenant les deux composants est préparée au moyen d'un procédé d'échange d'ions dans lequel une composition du matériau acide solide et des liants selon les revendications 21, 22 ou 23 est introduite dans une solution aqueuse contenant un sel de cuivre et éventuellement un sel d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, le mélange est laissé sous agitation pendant plusieurs heures, le solide en suspension est récupéré par filtration, lavé avec de l'eau déminéralisée et séché.
35. Procédé selon la revendication 24, dans lequel la zone contenant les deux composants est préparée au moyen de l'extrusion, du séchage et de la calcination d'un mélange mécanique des deux composants mélangés sous la forme d'une pâte avec un agent peptisant.
36. Procédé selon la revendication 24, dans lequel la zone contenant les deux composants est préparée au moyen de l'extrusion, du séchage et de la calcination d'un mélange mécanique du composant acide solide, du composant métallique et des liants selon les revendications 21, 22 ou 23.
37. Procédé selon la revendication 24, dans lequel la composition catalytique est constituée d'une zone unique contenant à la fois les composants métallique et acide solide.
38. Procédé selon la revendication 24, dans lequel la composition catalytique est constituée de deux zones distinctes ou plus, chacune contenant à la fois les composants métallique et acide solide, où la composition des zones uniques est différente.
39. Procédé selon les revendications 24 ou 25, dans lequel la composition catalytique est constituée de deux zones distinctes ou plus, chacune contenant le composant métallique, tandis que les autres contiennent une ou plusieurs combinaisons différentes des deux composants métallique et acide solide.
40. Procédé selon les revendications 24 ou 25, dans lequel la composition catalytique est constituée de deux zones distinctes ou plus, chacune contenant le composant métallique, tandis que les zones subséquentes contiennent le composant acide solide.
41. Procédé selon les revendications 24 ou 25, dans lequel la composition catalytique est constituée d'une zone unique contenant les deux composants de la composition catalytique, mélangés mécaniquement l'un avec l'autre.
42. Procédé selon les revendications 24 ou 25, dans lequel la composition catalytique est constituée de deux zones distinctes ou plus, chacune contenant le composant métallique, tandis que les autres contiennent des paires différentes des deux composants métallique et acide solide, mélangés mécaniquement.
43. Procédé selon les revendications 39, 40 ou 42, dans lequel la zone contenant le composant métallique est celle qui entre en premier en contact avec le courant des réactifs constitués du composant aromatique, de la cétone et de l'hydrogène.
44. Procédé selon la revendication 24, dans lequel la composition catalytique contient le composant métallique avec un gradient d'activité qui diminue dans une direction et le composant acide solide avec un gradient d'activité qui diminue dans la direction opposée.
45. Procédé selon la revendication 44, dans lequel le gradient d'activité du composant métallique diminue le long de la direction d'alimentation et de l'écoulement des réactifs, tandis que le gradient d'activité du composant acide solide diminue le long de la direction et de l'écoulement opposés.

46. Procédé selon les revendications 1 ou 2, réalisé à une température située dans la plage de 50 à 350 °C et à une pression égale ou supérieure à la pression atmosphérique.

47. Procédé selon la revendication 46, réalisé à une température située dans la plage de 100 à 250 °C et à une pression située dans la plage de 1 à 50 bars.

48. Procédé selon les revendications 1 ou 2, dans lequel un rapport molaire qui n'est pas inférieur à 1 est utilisé dans l'alimentation entre l'hydrocarbure aromatique et la cétone.

49. Procédé selon la revendication 48, dans lequel le rapport molaire est supérieur à 2.

50. Procédé selon les revendications 1 ou 2, dans lequel un rapport molaire qui n'est pas inférieur à 1 est utilisé entre l'hydrogène et la cétone dans l'alimentation.

51. Procédé selon la revendication 50, dans lequel un rapport molaire supérieur à 2 est utilisé entre l'hydrogène et la cétone dans l'alimentation.

52. Procédé selon l'une quelconque des revendications précédentes, dans lequel le composé aromatique est le benzène, la cétone est l'acétone, et le produit résultant du procédé d'alkylation, qui contient de l'isopropylbenzène, du benzène non converti, de l'hydrogène non converti, de l'eau et des polyisopropylbenzènes, est fractionné dans une section de séparation pour obtenir une première fraction contenant principalement de l'hydrogène, une deuxième fraction contenant principalement de l'eau, une troisième fraction contenant principalement du benzène, une quatrième fraction contenant principalement de l'isopropylbenzène et une cinquième fraction contenant principalement des polyisopropylbenzènes, où la première fraction est réutilisée dans l'étape de réaction avec l'acétone et le benzène, la deuxième fraction est prélevée du procédé, la troisième fraction est partiellement réutilisée dans l'étape de réaction avec l'acétone et l'hydrogène et partiellement dans une étape subséquente de transalkylation, où elle est mise à réagir avec la cinquième fraction pour produire de l'isopropylbenzène.

53. Procédé selon la revendication 52, dans lequel le produit de la réaction de transalkylation est fractionné en benzène, qui est rechargé dans le procédé d'alkylation, cumène et polyisopropylbenzènes, où les polyisopropylbenzènes sont rechargés dans l'étape de transalkylation.

54. Procédé de production de phénol qui comprend les étapes suivantes :

- (a) la réaction de benzène, d'acétone et d'hydrogène en présence d'un système catalytique comprenant un matériau acide solide, du cuivre, et éventuellement un ou plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides,
- (b) l'oxydation du cumène en hydroperoxyde de cumène,
- (c) le réarrangement de l'hydroperoxyde de cumène en phénol et acétone,

caractérisé en ce que l'étape (a) est effectuée selon une ou plusieurs des revendications 1 à 53.

55. Procédé selon la revendication 54, dans lequel l'acétone qui est formée dans l'étape (c) est recyclée dans l'étape de synthèse (a) de cumène.

56. Composition catalytique contenant une zéolite sous forme acide, Cu et un ou plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, à condition que lorsque la zéolite est la zéolite Y et la composition catalytique contient un ou plusieurs éléments du groupe VIB, ledit élément soit choisi entre Mo et W.

57. Composition catalytique selon la revendication 56, contenant une zéolite sous forme acide choisie parmi la zéolite beta, la zéolite ZSM-12 et la mordenite, Cu et un ou plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides.

58. Composition catalytique selon la revendication 56, contenant de la zéolite Y sous forme acide, Cu et un ou plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides.

59. Composition catalytique selon la revendication 56 contenant un liant.

60. Composition catalytique selon les revendications 56 ou 59, dans laquelle la zéolite est la zéolite beta.

5 61. Composition catalytique selon la revendication 60, contenant de la zéolite beta, du cuivre, un élément choisi parmi Cr et Al, et éventuellement un liant.

62. Composition catalytique selon la revendication 61, dans laquelle le cuivre et le chrome sont sous la forme de chromite de cuivre.

10 63. Procédé de préparation de la composition catalytique selon la revendication 56, comprenant l'imprégnation de la zéolite avec une solution aqueuse contenant un sel de cuivre et un sel d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, le séchage et la calcination.

15 64. Procédé selon la revendication 63, dans lequel le matériau résultant est lié avec un liant selon les revendications 21, 22 ou 23.

20 65. Procédé de préparation de la composition catalytique selon la revendication 56, comprenant l'imprégnation de la zéolite et des liants selon les revendications 21, 22 ou 23 avec une solution aqueuse contenant un sel de cuivre et un sel d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, le séchage et la calcination.

25 66. Procédé de préparation de la composition catalytique selon la revendication 56, comprenant l'échange ionique de la zéolite avec une solution aqueuse contenant un sel de cuivre et un sel d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, le séchage et la calcination.

30 67. Procédé de préparation de la composition catalytique selon la revendication 66, dans lequel le matériau obtenu est lié avec un liant.

35 68. Procédé de préparation de la composition catalytique selon la revendication 56, comprenant l'échange ionique d'une composition de zéolite et de liants appropriés avec une solution aqueuse contenant un sel de cuivre et un sel d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, le séchage et la calcination.

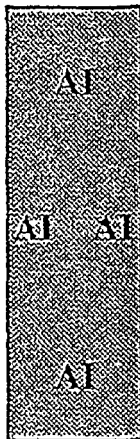
40 69. Procédé de préparation de la composition catalytique selon la revendication 56, comprenant l'extrusion, le séchage et la calcination d'un mélange mécanique, mélangé sous forme de pâte avec un agent peptisant, de zéolite, Cu et d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, sous la forme d'oxydes.

45 70. Procédé de préparation de la composition catalytique selon la revendication 56, comprenant l'extrusion, le séchage et la calcination d'un mélange mécanique, mélangé sous forme de pâte avec un agent peptisant, de zéolite, Cu et d'un ou de plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, sous la forme d'oxydes, et de liants selon les revendications 21, 22 ou 23.

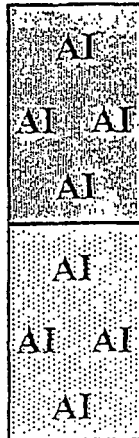
50 71. Procédé de préparation de la composition catalytique selon la revendication 56, qui comprend l'assemblage de diverses zones distinctes, chacune contenant le composant métallique contenant du cuivre et un ou plusieurs éléments choisis parmi des éléments des groupes IIIA, IVA, IIIB, IVB, VB, VIB, VIIB, du groupe VIII limité à Fe, Ru et Os, et de la série des lanthanides, ou le composant zéolitique, ou les deux composants.

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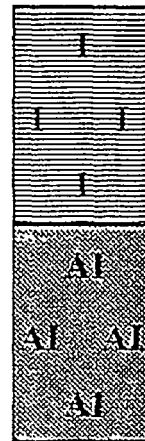
Fig. 1



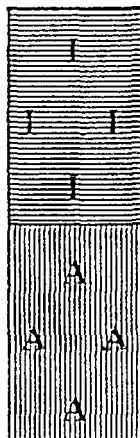
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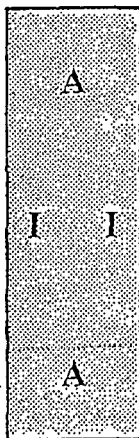
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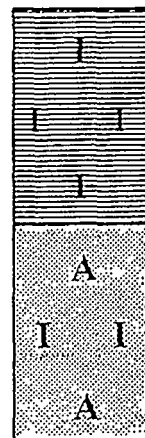
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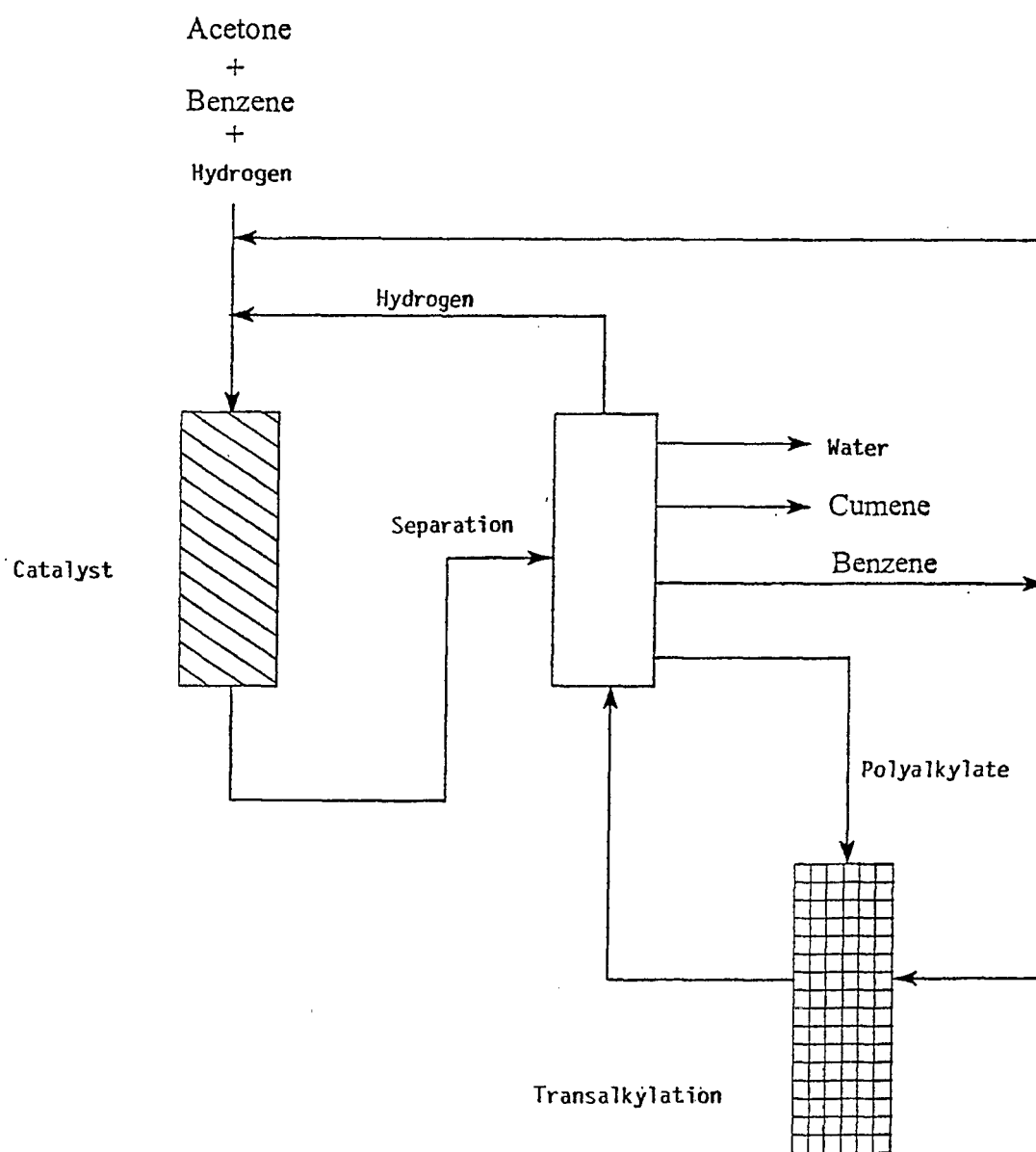


1e



1f

Fig. 2.



REFERENCES CITED IN THE DESCRIPTION

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