EUROPEAN PATENT SPECIFICATION

(54) Method for the separation of water from a reaction mixture using a hydroxy sodalite membrane
Verfahren zur Abtrennung von Wasser aus einem Reaktionsgemisch unter Verwendung einer Hydroxy-
Sodalit-Membran
Procédé pour la séparation d'eau d'un mélange réactionnel au moyen d'une membrane hydroxy sodalite

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Background of the invention

[0001] The present invention is directed to a process for performing a chemical reaction, to a process for separating water from a mixture containing water, to a membrane suitable for such processes and to a process for preparing a membrane.

[0002] Modern chemical engineering has a focus on the integration of reaction and separation. In-situ separation of water from the Fischer-Tropsch reaction mixture is an example of such integration. The continuous removal of water, which is produced as a side product, contributes to an improvement of conversion, product selectivity and catalyst lifetime.

[0003] More in general it can be remarked, that the removal of water from mixtures with water is a very important aspect in various areas of chemical engineering and process technology. In reactions where water is a by-product, be it an equilibrium reaction or a non-equilibrium reaction, the continuous removal of water produced, from the reaction mixture helps to improve the reaction efficiency.

[0004] Separating of water from mixtures thereof, more in particular with H₂, CO, CO₂, CH₄ and higher alkanes, is also very important, more in particular in case the other component(s) need(s) to be dry, for example for further reaction or use.

[0005] Separation can be based on several principles. With a hydrophilic membrane, it is possible to separate water from for example a mixture of water and an organic component. The separation principle is based on the preference of the membrane for adsorbing water. Also separation on size is possible, when the components to be separated are small enough in kinetic diameter to migrate through the zeolite pores and the components from which they have to be separated have a kinetic diameter that is too large.

[0006] The kinetic diameter can be understood as the diameter of a pore needed to let that specific molecule pass. Water has a kinetic diameter of 2.65 Å, even smaller than that of hydrogen. Thus separation based on geometry is possible, when one finds a zeolite with a pore size large enough to enable water to pass and to exclude hydrogen and other larger molecules. It was generally assumed that 6-membered rings were unsuitable for this type of separation.

[0007] In most chemical reactions with water as by-product, water has the smallest kinetic diameter. For example in a Fischer-Tropsch reaction mixture the kinetic diameters of H₂, CO and linear alkanes are respectively 2.89, 3.76 and 4.3 Å.

[0008] US-6,403,660 discloses a process for producing hydrocarbons. Water formed as a by-product is separated from the reaction medium using a zeolite membrane chosen from mordenite, ZSM-5, zeolite A and chabazite.

Summary of the invention

[0009] The invention is based on the surprising discovery that it is possible to continuously separate water from reaction mixtures, more in particular Fischer-Tropsch reaction mixtures, by the use of a hydroxy sodalite membrane.

[0010] Sodalite is a zeolite having as the largest apertures exclusively 6-membered rings, so-called sodalite cages. Hydroxy sodalite (sodalite hydrate, hydroxo sodalite) is the hydrated form of sodalite. It can be synthesized without a template from an aqueous solution and then contains water molecules in all its pores. The hydroxy sodalite is used without activation.

[0011] A definition of hydroxy sodalite can be found in "Atlas of Zeolite Frameworks, 5th revised Ed. Elsevier 2001, pages 254 and 255. In Figure 1, the structure has been given.

[0012] Hydroxy sodalite belongs to the group of feldspathoids and is a so-called cubic feldspathoid. It is entirely built of the so-called β-cages. These cages are truncated octahedrons (left structure of Fig.1). A truncated octahedron has 6 square faces and 8 hexagonal faces, which give it 36 edges and 24 vertices. A T (tetrahedral) atom (Si, Al) is located at each vertex. Oxygen atoms are located between the tetrahedral atoms (but not necessarily on the edge). So the edges are not meant to portray bonds, but merely the geometry of the polyhedron. In sodalite, this polyhedral arrangement can be distorted so that the exact arrangement of oxygen atoms is not regular.

[0013] In a first embodiment the invention is directed to a process for performing a chemical reaction in a reaction mixture, which reaction produces water as by-product, wherein the reaction mixture is in contact with a hydroxy sodalite membrane, through which water produced during the reaction is removed from the reaction mixture.

[0014] A second embodiment is directed to a process for separating water from a mixture thereof with at least compound from the group of H₂, CO, CO₂, CH₄ and higher alkanes, said process comprising passing the mixture over a hydroxy sodalite membrane and removing the water through the membrane.

[0015] A third embodiment concerns an especially suitable membrane or use in these processes, namely a tubular hydroxy sodalite membrane supported on a tubular porous ceramic support.

[0016] A 4th and final embodiment of the invention concerns a process for preparing a supported hydroxy sodalite membrane said process comprising providing a suitable porous membrane support material, preferably a ceramic support
material, providing a solution of a silicate and an aluminate in water and reacting the solution under hydrothermal conditions to produce a hydroxy sodalite layer on the surface of the porous material.

Detailed embodiments

[0017] As indicated above, in a first embodiment, the invention resides therein that water is removed continuously from the reaction mixture of a reaction wherein water is a by-product. By passing the reaction mixture along the surface of a hydroxy sodalite membrane, the water is removed from the reaction mixture. Preferably the temperature is at least 100°C. Above this temperature the structure of the sodalite is slightly distorted thereby easing the transfer of water through the membrane, which would otherwise be very difficult, as the diameter of the 6-membered cages is slightly smaller than the kinetic diameter of water. Preferably the temperature is between 150°C and 500°C. Above these temperatures the stability of the hydroxy sodalite decreases.

[0018] Generally, there are two separation methods with membranes: pervaporation and gas or vapour permeation. Pervaporation is the separation through a membrane with on the feed side a liquid mixture and an applied vacuum or lower pressure on the permeate side which makes the permeated components evaporate. With gas permeation both the feed mixture at the feed side and the separated components on the permeate side are in gas phase.

[0019] It may be advantageous to use increased pressure, as this increases the partial pressure of the reaction mixture, in case the water is present in the gaseous phase. Suitable pressures are between 0.5 bar (abs) to 100 bar (abs).

[0020] The membrane is preferably supported on a suitable porous support, such as a metal or, most preferred, a ceramic support, such as sintered α-alumina. Optionally, a mesoporous layer, such as a silica or titania layer, may be present between the support and the hydroxy sodalite layer. The thickness of the membrane is preferably between 1 and 60 μm, more preferred between 5 and 20 μm.

[0021] The membrane has to have a closed structure, i.e. there should be substantially no cracks or holes in the membrane, other than the pores of the sodalite structure.

[0022] In order to improve the flux, it is possible to flush the permeate side of the membrane with a dry gas.

[0023] Suitable reactions in the context of the present invention are the Fischer-Tropsch synthesis, condensation reactions and dehydration reactions.

[0024] Besides the well-known conventional production of fuels from oil, that is called refining, there is a 'gas to liquids' process (named the Fischer-Tropsch process). In this process carbon monoxide and hydrogen, together called the synthesis gas, are converted over an iron or cobalt catalyst (in fact it is the catalytic hydrogenation of carbon monoxide) into a mixture of hydrocarbons and water [1,2]. The basic general reaction is as follows:

\[ n\text{CO} + (2n+1)\text{H}_2 \rightarrow C_n\text{H}_{2n+2} + n\text{H}_2\text{O} \]

[0025] The hydrocarbons are synthesized by a chain growth process, with the length of the chain dependent on the catalyst selectivity and reaction conditions. The products formed primarily are normal paraffins, but α-olefins and alcohols are formed as well.

[0026] Suitable reaction conditions for FT reactions comprises pressures of between 5 and 50 bar(abs) and temperatures between 150°C and 500°C, temperatures of about 250°C (low temperature reaction) or 450°C (high temperature) being preferred.

[0027] Water that is formed can react with carbon monoxide to form hydrogen and carbon dioxide in the so-called water-gas-shift reaction, an unwanted side reaction:

\[ \text{CO} + \text{H}_2\text{O} \rightleftharpoons \text{CO}_2 + \text{H}_2 \]

[0028] In the early Fischer-Tropsch (FT) process methane reacts with air over a catalyst to create the synthesis gas, but this syngas can also be obtained from coal or even biomass gasification. Next to the main advantage of the Fischer-Tropsch process, which is the independence from oil, there is the advantage of making a cleaner product that, for example, contains no sulphur or aromatics.

[0029] Removal of water accordingly is advantageous for the reaction, not only because the subsequent reaction of CO and water is suppressed and the volume of the reactor is used more efficiently, but also because no ageing of the catalyst occurs. In general the same arguments apply to the other reactions, such as condensation reactions (which are generally equilibrium reactions, so that automatically the removal of water drives the reaction in the right direction) and dehydration reaction, for example the dehydration of hydroxy-group containing hydrocarbons to unsaturated hydrocarbons, more in particular alkanoil to alkane.

[0030] For the second embodiment, the separation of water from various mixtures with other compounds, generally
the same considerations apply with respect to the temperature, pressure and membrane structure.

[0031] More in particular the invention is suitable for drying methane and higher alkanes (C\textsubscript{2}-C\textsubscript{10}), and removing water from H\textsubscript{2}, CO and CO\textsubscript{2}.

[0032] The invention is further directed to the membrane itself and the process of making it. The membrane itself has already been described in relation to the other two embodiments.

[0033] The process for preparing a suitable membrane, comprises providing a suitable porous membrane support material, preferably a ceramic support material, such as \(\alpha\)-alumina, steel coated with titania and the like providing a solution of a silicate and an aluminate in water and reacting the solution under hydrothermal conditions to produce a hydroxy sodalite layer on the surface of the porous material. Hydrothermal conditions are preferably temperature of at least 125\(^\circ\)C (up to 500\(^\circ\)C) for a period of at least 2 hours, preferably under autogenous conditions. Once the required sodalite material has formed, the reaction is stopped and the hydroxy sodalite is washed. The membrane is then ready for use, as no activation is needed. Actually, the conventional activation procedure to burn away a template is detrimental to the material.

[0034] The supported membrane modules may subsequently be fitted into a membrane cell, with one or more tubular modules, in which the ends of the tubes are fitted with sealing means to make sure that the inside and outside of the membranes are only connected through the pores of the membrane. This gas-tight membrane sealing is thus very important for a good functioning of the membrane. For the best results epoxy-resin sealing or graphite ring sealing are preferred, since. (fluorelastomere) rubber rings failed.

[0035] The membranes of the invention have a flux which may be at least 1 kg of water/m\textsuperscript{2}h, even up to 2 kg/m\textsuperscript{2}h or higher. Based on these values the skilled person can easily calculate the membrane surface area required for a given reaction or separation. For example for a commercial FT reactor at industrial scale producing 1250 ton/day, a membrane surface of about 10,000 m\textsuperscript{2} would suffice.

Examples

Preparation of a membrane

[0036] \(\alpha\)-alumina tubes having the following dimensions: 21 mm outer diameter, 18 mm inner diameter, 58 mm in length and a wall thickness of 1.5 mm were prepared. The pore size distribution is symmetrical (radially) with average pore size 0.07 - 0.1 microns. The whole tube consists of \(\alpha\)-alumina, there is no \(\gamma\)-alumina present (Steenkamp, Ph.D. thesis, University Twente)

[0037] The tubes have been sintered from Sumitomo AKP alumina powders at 1050 \(^\circ\)C for 60 minutes, with a heating/cooling rate from and to room temperature of 1 \(^\circ\)C/min.

[0038] Prior to synthesis, the tubes need to be cleaned, in order to remove dust and other impurities. After the support was manufactured cleaning with an ammonium hydroxide and hydrogen peroxide solutions mixture was carried out.

[0039] Just before the synthesis the supports were washed with ethanol solutions, and finally with water. The latter two washings were done in sonificated baths. The outer side of the tubes was covered with a teflon tape (the one used for example to make tap water lines leak-proof). Ultimately runs have been performed with epoxy-resin sealing on Teflon holders.

[0040] Syntheses in rotating autoclaves seemed (according to SEM analysis) to give a better zeolite layer on the support. One explanation could be the removal of little air bubbles on the surface of the support by the continuous rotation. Defects in the zeolite layer can be accredited to these bubbles. The second modification was a change in nutrient concentration. SEM-pictures showed large agglomerates of crystals (so called ‘cauliflowers’) freely formed as well as attached to the support and zeolite layer. These agglomerates indicate a too large amount of nutrients, which induces unwanted free crystal growth next to the (wanted) zeolite crystal growth on the support. Several syntheses with decreased amounts of Si and Al (but always with a high pH) lead to a final modified synthesis mixture composition (table 1)

<table>
<thead>
<tr>
<th></th>
<th>weight [g]</th>
<th>Mw [g mol\textsuperscript{-1}]</th>
<th>mol</th>
<th>molar ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium hydroxide</td>
<td>3.495</td>
<td>40</td>
<td>0.087375</td>
<td>16</td>
</tr>
<tr>
<td>Sodium aluminate</td>
<td>0.223</td>
<td>84</td>
<td>0.002654</td>
<td>0.5</td>
</tr>
<tr>
<td>Sodium metasilicate</td>
<td>1.134</td>
<td>212</td>
<td>0.005349</td>
<td>1</td>
</tr>
<tr>
<td>Water</td>
<td>47.064</td>
<td>18</td>
<td>2.615</td>
<td>489</td>
</tr>
</tbody>
</table>

[0041] Prior to the syntheses two mixtures were made in two separate polyethylene flasks. One (mixture A, table 2)
contained the silica source, in this case sodium metasilicate dissolved in water with a high pH (so with a part of the sodium hydroxide). The other mixture (mixture B, table 3) also contained NaOH in water, and with the aluminium source, sodium aluminate. Both mixtures were well stirred for half an hour (in order to dissolve and hydrolyse the oxides) at ambient temperature.

<table>
<thead>
<tr>
<th>Components mixture A</th>
<th>weight [g]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium metasilicate</td>
<td>1.134</td>
</tr>
<tr>
<td>Sodium hydroxide</td>
<td>1.483</td>
</tr>
<tr>
<td>Water</td>
<td>23.532</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Components mixture B</th>
<th>weight [g]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium aluminate</td>
<td>0.223</td>
</tr>
<tr>
<td>Sodium hydroxide</td>
<td>2.012</td>
</tr>
<tr>
<td>Water</td>
<td>23.532</td>
</tr>
</tbody>
</table>

[0042] Subsequently the two mixtures were poured together and stirred well for another half hour (in order to obtain complete mixing of the components). In order to prevent pre-crystallisation half an hour aging was applied when preparing synthesis mixtures for zeolite growth on supports.

[0043] After this the mixture was put in a stainless steel autoclave with a teflon inner liner and the support (whether stainless steel with an TiO₂ layer or α-alumina). The autoclave was put in an oven (on a rotating axis in case of the rotating synthesis) for 3 hours at 140 °C. Afterwards the autoclave was well cooled with tap water. Not only to stop the crystallisation (if the reaction continues zeolite A will be formed), but also for safety reasons: the caustic solution inside the autoclave is at its autogenic pressure. Subsequently the zeolites were filtered from the mixture and washed with water. In case of on-support growth, the ‘coated’ support is cleaned thoroughly with water.

[0044] XRD provided clarification that sodalite was made and whether other crystalline phases were present (zeolite A, quartz). For powders a Philips PW 1840 was used. Supported zeolite layers were identified by the Philips XRD of the Department of Materials Science of DelftChemTech (DCT), where also some sodalite powders have been analysed. Figure 2

[0045] SEM pictures were taken on a Philips XL20 SEM at the Polymers Department (PME) of Delft Technical University and at a Philips SEM with EDAX elemental analysis at the TNO Prins Maurits Laboratories (PML) in Rijswijk. Figure 3

[0046] Tests were carried out at temperatures around 160 °C and low pressures. The pressure should be high enough to have a certain pressure drop over the membrane. A pressure drop of 2 bar was chosen and with the permeate side at atmospheric pressure, a total absolute pressure of 3 bar at the feed side was used. To ensure that the mixture was in the vapour phase, temperatures were chosen well enough above the boiling temperatures of octane and water at 3 bara. A working temperature was chosen well above the boiling point of least volatile component (i.e. octane) at the specific pressure applied.

[0047] The resulting permeate was obtained and analysed. No Octane was found therein, only water.

Claims

1. Process for performing a chemical reaction in a reaction mixture, which reaction produces water as by-product, wherein the reaction mixture is in contact with a hydroxy sodalite membrane, through which water produced during the reaction is removed from the reaction mixture.

2. Process according to claim 1, wherein the said hydroxy sodalite membrane is supported on a porous support, preferably a ceramic support.
3. Process according to claim 2, wherein the support is an α-alumina support.

4. Process according to claim 1-3, wherein the removal of water through the membrane occurs at elevated temperature, preferably at least 100°C, more preferably between 150°C and 500°C.

5. Process according to claim 1-5, wherein the said chemical reaction is selected from the group of Fischer-Tropsch reactions, condensation reactions and dehydration reactions.

6. Process according to claim 5, wherein the said reaction comprises a Fischer-Tropsch synthesis or the production of alkene from alkanol.

7. Process according to claim 1-6, wherein the reaction is a slurry phase reaction and the reaction mixture is continuously passed along the surface of the membrane, the membrane being a tubular supported membrane.

8. Process for separating water from a mixture thereof with at least compound from the group of H₂, CO, CO₂, CH₄ and higher alkanes, said process comprising passing the mixture over a hydroxy sodalite membrane and removing the water through the membrane.

9. Process according to claim 8, wherein the said hydroxy sodalite membrane is supported on a porous support, preferably a ceramic support.

10. Process according to claim 9, wherein the support is an α-alumina support.

11. Process according to claim 8-10, wherein the removal of water through the membrane occurs at elevated temperature, preferably at least 100°C, more preferably between 150°C and 500°C.

12. Process according to claim 8-11, wherein the said membrane is a tubular membrane.

13. Tubular hydroxy sodalite membrane supported on a tubular porous ceramic support.

14. Membrane according to claim 13, wherein the support is an α-alumina support.

15. Membrane according to claim 13 or 14, wherein between the hydroxy sodalite membrane and the ceramic support, an intermediate layer of mesoporous material, such as a silica, is present.

16. Process for preparing a supported hydroxy sodalite membrane, suitable for use in the processes of claims 1-12, said process comprising providing a suitable porous membrane support material, preferably a ceramic support material, providing a solution of a silicate and an aluminate in water and reacting the solution under hydrothermal conditions to produce a hydroxy sodalite layer on the surface of the porous material.

17. Process according to claim 16, wherein the hydrothermal reaction is stopped once the membrane has been formed.

18. Process according to claim 16 or 17, wherein the said hydrothermal conditions comprise a temperature of at least 125°C for a period of at least 2 hours under an autogenous or higher pressure.

**Patentansprüche**

1. Verfahren zur Durchführung einer chemischen Reaktion in einem Reaktionsgemisch, wobei die Reaktion Wasser als Nebenprodukt erzeugt, wobei die Reaktionsmischung in Kontakt mit einer Hydroxy-Sodalith-Membran steht, durch welche das während der Reaktion entstehende Wasser aus dem Reaktionsgemisch entfernt wird.

2. Verfahren nach Anspruch 1, wobei die Hydroxy-Sodalith-Membran durch einen porösen Träger gestützt wird, vorzugsweise einen keramischen Träger.

3. Verfahren nach Anspruch 2, wobei der Träger ein α-Aluminiumoxidträger ist.

4. Verfahren nach Anspruch 1 bis 3, wobei die Entfernung von Wasser durch die Membran bei einer erhöhten Tem-
5. Verfahren nach Anspruch 1 bis 5, wobei die chemische Reaktion aus der Gruppe der Fischer-Tropsch-Reaktionen, der Kondensationsreaktionen und der Dehydrationsreaktionen ausgewählt ist.


7. Verfahren nach Anspruch 1 bis 6, wobei die Reaktion eine Aufschlammreaktion ist und die Reaktionsmischung kontinuierlich entlang der Oberfläche der Membran geleitet wird, wobei die Membran eine röhrförmige, gestützte Membran ist.

8. Verfahren zum Abdrengen von Wasser aus einer Mischung desselben mit wenigstens einer Verbindung aus der Gruppe H₂, CO, CO₂, CH₄ und höherer Alkane, wobei das Verfahren beinhaltet, dass die Mischung über eine Hydroxy-Sodalith-Membran geleitet wird und das Wasser durch die Membran entfernt wird.

9. Verfahren nach Anspruch 8, wobei die Hydroxy-Sodalith-Membran durch einen porösen Träger gestützt wird, vorzugsweise einen keramischen Träger.

10. Verfahren nach Anspruch 9, wobei der Träger ein α-Aluminiumoxidträger ist.

11. Verfahren nach Anspruch 8 bis 10, wobei die Entfernung von Wasser durch die Membran bei einer erhöhten Temperatur erfolgt, vorzugsweise zumindest 100°C, weiter bevorzugt zwischen 150°C und 500°C.

12. Verfahren nach Anspruch 8 bis 11, wobei die Membran eine röhrförmige Membran ist.

13. Röhrförmige Hydroxy-Sodalith-Membran gestützt durch einen röhrförmigen, porösen keramischen Träger.


17. Verfahren nach Anspruch 16, wobei die hydrothermale Reaktion gestoppt wird, sobald die Membran gebildet worden ist.

18. Verfahren nach Anspruch 16 oder 17, wobei die hydrothermalen Bedingungen eine Temperatur von zumindest 125°C für einen Zeitraum von wenigstens zwei Stunden unter einem autogenen oder höheren Druck aufweisen.

Reivendications

1. Procédé pour mettre en oeuvre une réaction chimique dans un mélange réactionnel, ladite réaction produisant de l'eau comme sous-produit, dans lequel le mélange réactionnel est en contact avec une membrane d'hydroxy sodalite, par l'intermédiaire de laquelle l'eau produite lors de la réaction est retirée du mélange réactionnel.

2. Procédé selon la revendication 1, dans lequel ladite membrane d'hydroxy sodalite repose sur un support poreux, de préférence un support en céramique.

3. Procédé selon la revendication 2, dans lequel le support est un support α-alumine.

4. Procédé selon les revendications 1-3, dans lequel le retrait de l'eau par l'intermédiaire de la membrane se produit
à température élevée, de préférence au moins 100°C, plus préférentiellement comprise entre 150°C et 500°C.

5. Procédé selon les revendications 1-5, dans lequel ladite réaction chimique est choisie parmi le groupe des réactions de Fischer-Tropsch, des réactions de condensation et des réactions de déshydratation.

6. Procédé selon la revendication 5, dans lequel ladite réaction comprend une synthèse de Fischer-Tropsch ou la production d'alcool à partir d'alcool.

7. Procédé selon les revendications 1-6, dans lequel la réaction est une réaction de type phase « slurry » et le mélange réactionnel est passé en continu le long de la surface de la membrane, la membrane étant une membrane tubulaire reposant sur un support.

8. Procédé de séparation de l'eau d'un mélange d'au moins un composé du groupe formé par H₂, CO, CO₂, CH₄ et d' alcanes supérieurs, ledit procédé comprenant le passage du mélange sur une membrane d'hydroxy sodalite et le retrait de l'eau par l'intermédiaire de la membrane.

9. Procédé selon la revendication 8, dans lequel ladite membrane d'hydroxy sodalite repose sur un support poreux, de préférence un support en céramique.

10. Procédé selon la revendication 9, dans lequel le support est un support α-alumine.

11. Procédé selon les revendications 8-10, dans lequel le retrait de l'eau par l'intermédiaire de la membrane se produit à température élevée, de préférence au moins 100°C, plus préférentiellement entre 150°C et 500°C.

12. Procédé selon les revendications 8-11, dans lequel ladite membrane est une membrane tubulaire.

13. Membrane tubulaire d'hydroxy sodalite reposant sur un support tubulaire en céramique poreux.

14. Membrane selon la revendication 13, dans laquelle le support est un support α-alumine.

15. Membrane selon les revendications 13 ou 14, dans laquelle une couche intermédiaire d'un matériau mésoporeux, telle qu'une silice, est disposée entre la membrane d'hydroxy sodalite et le support en céramique.

16. Procédé de préparation d'une membrane d'hydroxy sodalite reposant sur un support, ladite membrane étant appropriée pour une utilisation dans les procédés des revendications 1-12, ledit procédé comprenant la fourniture d'un matériau de support poreux adapté à la membrane, de préférence un matériau de support en céramique, la fourniture d'une solution d'un silicate et d'un aluminate dans de l'eau et la réaction de la solution dans des conditions hydrothermales produisant une couche d'hydroxy sodalite sur la surface du matériau poreux.

17. Procédé selon la revendication 16, dans lequel la réaction hydrothermale est interrompue lorsque la membrane est formée.

18. Procédé selon les revendications 16 ou 17, dans lequel lesdites conditions hydrothermales comprennent une température d'au moins 125°C appliquée sur une période d'au moins 2 heures sous une pression autogène ou supérieure.
SEM pictures taken at the Philips XL Scanning Electron Microscope at the TNO PM Laboratories.

Figure 3
REFERENCES CITED IN THE DESCRIPTION

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Patent documents cited in the description

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Non-patent literature cited in the description