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Description

[0001] The present invention relates to an advanced method for removing suspended organic material such as tar from gas flows. Such gas flows may originate i.a. from gasification of biomass, organic waste or coal.

Background

[0002] A method for clarifying gas streams is known from WO 2008/010717. In this method, gas leaving a biomass reactor is subjected to a hydrocarbon oil flow. Excess oil is recirculated and is used after filtration for contact with the gas from the reactor. Additional oil is received from a downstream separation device into which the gas is entered.

[0003] WO 03/018723 discloses a so-called OLGA system for gasifying biomass. The gas which results from gasification is subjected to a two-step cleaning treatment for removing tars. In a first step the gas is condensed in a first cleaning fluid which is a hydrocarbon oil. Saturation can take place for example by spraying oil in the gas stream. In a second stage oil is used for absorption of the remaining tars in an absorption column. After use the oil with tars is discharged to a separator wherein the heavy fractions are returned to the biomass gasifier and the lighter fractions are further used as oil for the above process.

[0004] WO 2011/037463 discloses an oil recovery system (ORS) for a more effective tar removal from gasified biomass. It uses a first cleaning oil which is based on aromatic hydrocarbons and wherein the mixture of tar and first cleaning oil is separated into a light fraction and a heavy fraction, and the light fraction is reused as first cleaning oil. The second cleaning oil for removing residual tar components is based on aliphatic hydrocarbons.

[0005] The use of hydrocarbon oils for removing tars has some disadvantages, such as a limited stability at high temperatures and in the presence of water and oxidative agents, resulting in degradation and loss of oil due to the significant volatility of the hydrocarbons. As a result, the scrubbing liquid and equipment are contaminated and the process must be interrupted at regular intervals for exchange of scrubbing liquid and cleaning of equipment.

[0006] US 5772734 discloses the use of a wide variety of organic scrubbing liquids, including low-viscosity silicone oils, light minerals oils and glycol ether especially triethylene glycol dibutyl ether for removing organic compounds such as chlorinated hydrocarbons or aromatic hydrocarbons (such as toluene) from industrial gas streams.

[0007] It was found that the problems associated with prior art processes can be effectively solved by using a scrubbing oil based on a polysiloxane.

Description of the invention

[0008] The invention thus pertains to a process of clarifying a gas stream according to claim 1. The contacting (scrubbing) results in absorption of tar-like compounds, namely (poly)aromatic compounds of more than 6 carbon atom, especially 9-18 carbon atoms, into the polysiloxane scrubbing liquid. The gas stream to be clarified can be any gas comprising small molecules such as hydrogen, nitrogen, small hydrocarbons (up to 4 carbon atoms, such as methane and ethane), carbon monoxide and carbon dioxide. For example, the gas may comprise hydrogen, carbon monoxide, carbon dioxide, methane, nitrogen, and water (vapour), in addition to the tar components. The gas is in particular an energy gas or a synthesis gas containing one or more of hydrogen, carbon monoxide, and methane, in addition to varying levels of nitrogen, carbon dioxide and/or water, and sometimes ammonia and small hydrocarbons other than methane.

[0009] More in particular, the gas comprises at least 30 vol.%, especially more than 40% of one or more of hydrogen, methane and carbon monoxide. A typical composition of a gasification mixture comprises 10-40% CO, 5-30% CO₂, 2-40% H₂, 4-24% CH₄ and 5-40% H₂O (based on volumes). Tar-like components may be present in widely varying concentrations, e.g. from 1 ppm to 2% or higher, of which benzene may be the most abundant one (e.g. up to 1% (v/v) and up to 10% of the energy of the gasification mixture). In case of gasification using air, these levels are roughly divided by 2, the remaining 40-60% of the mixture being nitrogen.

[0010] Such a gas may originate from the gasification of biomass, organic waste, coal or a mixture of these. The organic waste may e.g. be municipal solid waste (MSW) and materials resulting in refuse-derived fuel (RDF). The biomass may be agricultural or forestry residues (wood chips, leaves, straw, grass, etc.) and the like.

[0011] The organic polysiloxane to be used as a washing (scrubbing) liquid comprises an alternating silicon-oxygen chain wherein the silicon is further substituted with organic groups. The polysiloxane is an aryl polysiloxane, i.e. at least a part of the organic groups are aryl groups, including aralkyl or alkaryl groups. Preferably, the polysiloxane comprises an average of between 0.2 and 1.8 C₅-C₁₄ aryl group, more preferably 0.8-1.8 C₅-C₁₀ aryl group, per silicon atom. Advantageously, the polysiloxane also comprises alkyl groups, in particular 0.2-1.8 C₁-C₆ alkyl group per silicon atom, more in particular 0.2-1.8 C₁-C₄ alkyl group per silicon atom.

[0012] In a preferred embodiment, the polysiloxane comprises an average of between 0.5 and 1.5 C₅-C₁₀, more preferably 0.5-1.5 C₅-C₁₀ aryl group, most preferably 0.5-1.5 C₆-C₈ aryl group per silicon atom and/or between 0.5 and 1.5 C₁-C₄ alkyl group per silicon atom. The polysiloxane preferentially has a molar weight between 500 and 14,000.
Da, preferably between 700 and 7,000 Da, more preferably between 1,000 and 5,500.

[0013] The polysiloxane can be represented by one of the formulas (I) and (II):

(I) \[-\text{Si}^{\text{R1}}\text{R2}-\text{O}-\text{n}\]  

(II) \[-\text{Si}(\text{R1})_{2}-\text{O}-\text{n1}[-\text{Si}(\text{R2})_{2}-\text{O}-\text{n2}[-\text{Si}^{\text{R1}}\text{R2}-\text{O}-\text{n3}\]  

wherein:

- R1 and R2 are the same or different, optionally substituted C1-C14 hydrocarbyl groups,
- n is between 5 and 100, preferably between 7 and 40,
- n1 + n2 + n3 = n, (n1+n3)/n2 is between 1/9 and 9/1, preferably between 1/4 and 4/1, most preferably between 2/3 and 3/2, and/or (n2+n3)/n1 is between 1/9 and 9/1, preferably between 1/3 and 3/1, most preferably between 2/3 and 3/2.

[0014] Preferred polysiloxanes of formulas (I) and (II) are those wherein R1 comprises at least 50% (by number) of aryl groups, preferably of 5-14 carbon atoms, more preferably of 5-10 carbon atoms, most preferably of 6-8 carbon atoms, the remainder being e.g. alkyl groups. Preferably at least 80%, more preferably at least 95% of the groups R1 are such aryl groups. Aryl groups as used herein include alkylaryl and arylalkyl (aralkyl) groups. Suitable aryl groups include phenyl, methylphenyl (p-tolyl), methoxyphenyl, benzyl, 2-phenyl-isopropyl, naphthyl and the like. Preferably R2 comprises C1-C4 alkyl groups. Suitable alkyl groups include e.g. methyl and ethyl.

[0015] The polysiloxane can be a homopolymer, as represented by formula (I), or an alternating, block or random copolymer as depicted by formula (II), of dialkyl, alkyl-aryl and/or diaryl siloxanes. It can also be a mixture of different poly-dialkyl, polyaryl-alkyl and/or polydiaryl siloxanes. Preferably the polysiloxane is a poly(methylphenyl-siloxane) or a poly-diphenyl-co-dimethyl-siloxane. Preferably the proportion of aryl groups is between 0.5 and 1.5 aryl (e.g. phenyl) groups per silicon atom.

[0016] In the process of the invention, the gas is contacted with the polysiloxane at elevated temperatures, such as a temperature between 30 and 150 °C, preferably between 60 and 120 °C, to absorb therein the aromatic tar-like components. Contrary to prior art processes using hydrocarbon-based scrubbing liquids, the temperature need not be above the water dew point, since water can be absorbed into the liquid without causing problems. The possibility of operating below the water dew point constitutes another advantage of the process of the invention.

[0017] The above contacting (scrubbing) temperatures will apply at atmospheric or slightly superatmospheric conditions. When using higher pressures, the temperatures will typically be higher so as to have comparable vapour pressures of the various components. As a rough rule of thumb, a doubling of the pressure corresponds to a higher temperature of about 20°C. Thus, an absorption step performed at 80°C and 1 bar is roughly equivalent to a step performed at 100°C and 2 bar, or at 145°C at 10 bar, etc.

[0018] After being contacted with the gas, the polysiloxane, with the aromatic compounds absorbed therein, is heated to a temperature which is at least 50 °C, preferably between 80 and 120°C above the contacting temperature, and is stripped (desorbed) with a stripping gas, such as nitrogen, to remove the aromatic compounds. After stripping, the stripped polysiloxane is returned for a further contact cycle.

[0019] Again, these temperature differences apply at equal pressures. Instead of increasing the temperature, the pressure can be lowered, where, again, a temperature increase of 20°C arbitrarily corresponds to a pressure decrease of a factor 2 and a temperature increase of at least 50°C corresponds to a pressure decrease of a factor of at least 5.6. So, at equal temperatures, the stripping is preferably performed at a pressure which is a factor between 16 and 80 below the scrubbing pressure.

[0020] In absolute terms, the stripping can be performed for example at a temperature between 120 and 250 °C, in particular between 150 and 220 °C at atmospheric pressure. Alternatively, the preferred stripping temperatures are e.g. between 90 and 220°C, preferably between 120 and 190°C at 1/8 (0.125) bar.

[0021] The gas stream is to be cleared from heavier compounds which hinder the subsequent use of the gas stream. Compounds to be removed are tar-like or tar-constituting compounds, namely aromatic hydrocarbons having 6 or more carbon atoms, such as benzene, alkylbenzenes, naphthalene and alkynaphthalenes, and higher homologues, as well as the corresponding hydroxyl (phenolic) and amino compounds. The process of the invention is particularly useful for removing bicyclic, tricyclic and tetracyclic aromatic compounds having 9-18 carbon atoms, which are among the lighter tar components. These are largely carbocyclic, but heterocyclic compounds such as indoles, (iso)quinolines and
Examples of (poly)aromatic compounds which are effectively removed by the present process include indene, naphthalene, quinoline, isoquinoline, 2-methyl-naphthalene, 1-methyl-naphthalene, biphenyl, ethenyl-naphthalene, acenaphthylene, acenaphthene, fluorene, anthracene, phenanthrene, fluoranthene, pyrene, acenanthylene, benzoanthracenes and chrysene. In addition to the C9-C18 aromatic components, slightly lighter, e.g. having 7 or 8 carbon atoms, as well as heavier tar-like compounds, e.g. up to 24 carbon atoms are effectively removed by the present process as well. Examples include the lighter components phenol, cresols, xylenes, ethylbenzene, styrene, cumene and the like, as well as the heavier benzo-fluoranthenes, benzopyrenes, pyrene, picene, benzoperylenes, indenoperylenes, dibenzanthracenes, benzoperylenes, coronene etc.

It is preferred that if the gas stream contains substantial levels of tar-like components having more than 18 carbon atoms, in particular if it contains components of more than 24 carbon atoms, a pre-cleaning steps is incorporated in the clarification process, as further described below.

In an embodiment of the invention, when the gas contains substantial levels of heavier tars, the gas to be clarified is pre-treated with another scrubbing liquid prior to being contacted to the polysiloxane. Such a prior scrubbing liquid preferably has a high affinity to more complex polycyclic (aromatic) hydrocarbons. Suitable prior scrubbing liquids are aromatic hydrocarbons, for example polyaromatic compounds corresponding to the tar-like components present in the gas and/or originate from scrubbing the tar-containing gas, comprising polyaromatics having 2-4 rings. Commercially available coal tars or coal tar naphthas (the lighter equivalent of coal tar) mainly consisting of C7-C18 hydrocarbons, are suitable as starting material for prior scrubbing. They will be suppleted during scrubbing with tars extracted from the gas. The prior scrubbing can be performed at a higher temperature than the scrubbing with the polysiloxane, for example a liquid inlet temperature of between 150 and 300°C, and a gas inlet temperature between 250°C and 900 °C. Alternatively, or additionally, the gas issuing from the gasification may be cleared from dust particles before and/or after an optional prior scrubbing step by passing it e.g. through an electrostatic filter or an electrostatic precipitator as described in WO2008/010717. Other methods or devices for clearing dust and/or tar particles prior to the scrubbing step, can be used as well, such as an aerosol scavenger as described in WO 2011/099850.

In the prior scrubbing step using (poly)aromatic hydrocarbons as absorption liquid, the spent liquid resulting from the absorption step can be subjected to a separation step using evaporation of the lighter components. These lighter components can be reused as absorption (scrubbing) liquid for the prior scrubbing step. The heavier fraction resulting from the separation step is discharged. Part of this discharged heavy fraction can be returned to the inlet of the gasification or pyrolysis reactor and converted to lighter components, or it can be used for other purposes such as heating. Further details for a prior absorption (scrubbing) step using aromatic hydrocarbons are described in WO 2011/037463.

If desired, a further cleaning step can be introduced before or after the polysiloxane cleaning step, for example for further removing polar compounds such as ammonia or the like, using neutral, acidic or alkaline aqueous scrubbing liquids, or for removing water, e.g. by condensation. However, appreciable levels of water, ammonia, amines and the like are effectively captured by the polysiloxane liquid, so that additional cleaning steps will often not be necessary, unless the gas contains very high levels of such polar compounds. As these compounds such as water and ammonia are also readily desorbed from the polysiloxane liquid, leaving the scrubbing liquid clean and stable, this constitutes a major advantage of the process of the invention.

The operation of the process and system of the invention can be described with reference to the accompanying figure. The system comprises an absorbing unit 32 such as a column, in which the gas to be clarified is fed in one part, preferably the bottom part, through gas inlet 34 and the scrubbing liquid comprising the polysiloxane is introduced through liquid inlet 35 in another part, preferably in the top part. Contact between the up-flowing gas and the down-flowing liquid can be enhanced by conventional means such as by spraying, using a packed column or a plate column. Although concurrent scrubbing is feasible, a counter-current mode, using a packed column, is preferred. Clean gas can leave the absorbing unit at the top through 37, preferably after passing a mist-collecting unit so as to minimise loss of polysiloxane liquid. Spent scrubbing liquid in which the tar components are absorbed, is collected at the bottom and discharged through outlet 38. The absorbing unit can be operated at temperatures of e.g. between 30 and 150°C, at atmospheric or slightly superatmospheric pressures, or at higher temperatures, when higher pressures are applied.

Desorption of the spent scrubbing liquid is preferably performed in a stripping unit 46, where the tar-like components are desorbed from the polysiloxane scrubbing liquid by a stripping gas. The stripping unit can be a tray tower, packed column, bubble column, spray tower or the like. The stripping gas can be co-currently or, preferably, counter-currently contacted with the spent polysiloxane. The spent liquid is fed to the stripping unit through line 41, optionally containing a pump 40, a safety filter 42 and a heater 43, and introduced through inlet 48. The stripping gas is introduced at inlet 50 and the spent stripping gas is discharged through outlet 51. The stripping gas can e.g. be air, nitrogen, carbon dioxide or mixtures thereof. The stripping unit is operated at about 100°C above the temperature of the absorption column, more generally between 70 and 120°C above the temperature of the absorption column, when using the same...
pressures. At atmospheric pressures, the temperatures can be between 120 and 250°C. Instead of using higher temperatures, the stripping unit 46 can be operated at lower pressure than the absorption unit 32. The desorbed polysiloxane exits the stripper through 52 and is returned to the absorption column through line 54, optionally using through a pump 58 and a cooler 56. Heater 43 and cooler 56 can advantageously be in heat exchanging communication. Instead or in addition to the heater 43 and cooler 56, pressurising and depressurising devices can be inserted. The gas issuing from the stripping unit through 51 can be cooled for condensing and separating the tar components. If desired, these separated tar components can be used to adjust the content of aromatic hydrocarbons and/or the viscosity of upstream scrubbing liquids.

[0030] The system of the invention for producing and cleaning an energy gas or synthesis gas can comprise a coal, waste or biomass gasifier of pyrolyser 10 to which a flow of biomass is added through an inlet. The gasifying gas can be e.g. air, oxygen and/or steam. The biomass, coal and/or waste can be further supplied with tar-like components issuing for one of the downstream clarifying process steps. Gasification can be performed at a temperature of 600-1300°C using sub-stoichiometric quantities of oxygen. Pyrolysis can be performed at the same or somewhat lower temperature (e.g. from 450°C up to 950°C).

[0031] Gas issuing from the gasifier can be subjected to a first separation step 15 based on gravitation and more particularly with a cyclone for removing dust. Next, the partly cleaned gas flow can be treated in the prior scrubbing step 20. Part or most of the tars contained in the gas flow are caught in this way. Besides tars based on hydrocarbons and dust, also sulphur and chloride-containing material can be removed from the gas flow. In an optional next step, the partly cleaned gas can be passed through a filter 25, for example an electrostatic precipitator, which removes dust. Instead of scrubber 20 and/or filter 25, an alternative tar and dust removing step can be inserted, such as an aerosol scavenger. Then, the gas can be entered in the process of the invention through line 26 for removing residual tar using the polysiloxane scrubbing and enters the scrubber through inlet 34. The preceding units 10, 15, 20 and 25 are only schematically depicted in the accompanying figure. More details are given in WO 2008/010717 and WO 2011/037463. Depending on the quality of the input gas, one or more or all of the intermittent cleaning steps described above can be dispensed with.

**Description of the figure**

[0032] The accompanying figure shows a gas cleaning system according to the invention.

**Examples**

[0033] A substitute natural gas (SNG) containing 13 vol% methane, 2 vol% nitrogen, 32 vol% carbon monoxide, 18 vol% carbon dioxide, 28% vol% hydrogen and 4 vol% ethene and containing about 11-12 g/Nm³ of C8-C16 (poly)aromatic hydrocarbons, was subjected to absorption using either a conventional aliphatic hydrocarbon derived from mineral oil or a commercial polymethylphenylsiloxane (PMPS) as an absorption liquid, and the absorption liquid was subsequently stripped using air.

[0034] The setting of the absorber and stripper were as follows in both cases:

- Temperature absorber = 80°C
- Temperature stripper = 180°C
- Gas flow stripper = 16 l/min (~1 Nm³/h)
- Oil flow absorber/stripper = 2.1 l/min

[0035] The tests were performed 4 times and the results were averaged. Table 1 below shows the concentrations (in mg/Nm³) and removal rates for the hydrocarbons up to pyrene. Benzene and toluene were omitted as they could not be precisely measured. Tar components of a level below 50 mg/Nm³ (in both columns) were also omitted.

**Table 1: Removal rate of tar components by aliphatic oil vs. polysiloxane oil**

<table>
<thead>
<tr>
<th>Tar Component</th>
<th>Mineral Oil</th>
<th>PMPS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Abs in</td>
<td>Abs out</td>
</tr>
<tr>
<td>Ethylbenzene</td>
<td>14</td>
<td>2</td>
</tr>
</tbody>
</table>
Claims

1. A process of clarifying a gas stream containing tar-like or tar-constituting components comprising aromatic compounds having 6 or more carbon atoms, comprising contacting the gas with a liquid organic aryl polysiloxane as a washing liquid.

2. A process according to claim 1, wherein the polysiloxane comprises an average of between 0.2 and 1.8 C5-C14 aryl group per silicon atom.

3. A process according to claim 1 or 2, wherein the polysiloxane comprises an average of between 0.5 and 1.5 C5-C10 aryl group and between 0.5 and 1.5 C1-C4 alkyl group per silicon atom.

4. A process according to any one of claims 1-3, wherein the polysiloxane has a molar weight between 700 and 7000 Da.

5. A process according to any one of claims 1-3, wherein the polysiloxane is a polymethylphenylsiloxane or a polydiphenyl-dimethyl-siloxane.

6. A process according to any one of the preceding claims, wherein the gas comprises one or more of hydrogen, carbon monoxide, carbon dioxide, methane and nitrogen, in particular at least 30 vol.% of one or more of hydrogen, methane and carbon monoxide.

7. A process according to any one of the preceding claims, wherein the gas is to be clarified from tar-like components comprising polycyclic aromatic compounds having 9-18 carbon atoms.

8. A process according to any one of the preceding claims, wherein the gas is contacted with the polysiloxane at a contacting temperature between 30 and 150 °C, preferably between 60 and 120 °C, at atmospheric contacting pressure, or at a corresponding higher temperature at higher pressure.

9. A process according to claim 8, wherein the polysiloxane, after the contacting with the gas, is heated to a temperature which is higher than the temperature of the contacting step and/or is depressurised to a pressure which is lower than the contacting pressure, preferably in the presence of a stripping gas, and is subsequently returned for a further contact cycle.

10. A process according to claim 9, wherein, at equal pressure, the higher temperature is at least 50 °C above the contacting temperature, preferably between 80 and 120°C above the contacting temperature, or wherein, at equal
temperature, the lower pressure is at least 5.6 times, preferably between 8 and 16 times lower than the contacting pressure.

11. A process according to any one of the preceding claims, wherein the gas, prior to contacting to the polysiloxane, is treated with a scrubbing liquid comprising aromatic hydrocarbons, at a temperature of between 150 and 900 °C.

12. A process according to any one of the preceding claims, wherein the gas, prior to contacting to the polysiloxane, is subjected to an electrostatic filter and/or an aerosol scavenger.

13. A process according to any one of the preceding claims, wherein the gas originates from the gasification of biomass, organic waste, coal or a combination thereof.

Patentansprüche

1. Verfahren zur Reinigung eines Gasstroms, der teerartige oder teerhaltige Komponenten enthält, die aromatische Verbindungen mit 6 oder mehr Kohlenstoffatomen umfassen, bei welchem Verfahren das Gas mit flüssigem organischen Arylpolysiloxan als Waschflüssigkeit in Kontakt gebracht wird.

2. Verfahren nach Anspruch 1, bei dem das Polysiloxan durchschnittlich zwischen 0,2 und 1,8 C5- bis C14-Arylgruppen pro Siliciumatom umfasst.

3. Verfahren nach Anspruch 1 oder 2, bei dem das Polysiloxan durchschnittlich zwischen 0,5 und 1,5 C5- bis C10-Arylgruppen und zwischen 0,5 und 1,5 C1- bis C4-Alkylgruppen pro Siliciumatom umfasst.

4. Verfahren nach einem der Ansprüche 1-3, bei dem das Poly-siloxan ein molares Gewicht zwischen 700 und 7000 Da aufweist.

5. Verfahren nach einem der Ansprüche 1-3, bei dem das Polysiloxan ein Polymethylphenylsiloxan oder ein Polydi phenyldimethylsiloxan ist.


7. Verfahren nach einem der vorhergehenden Ansprüche, bei dem das Gas von teerartigen Komponenten gereinigt werden soll, die polycyclische aromatische Verbindungen mit 9 bis 18 Kohlenstoffatomen umfassen.


10. Verfahren nach Anspruch 9, bei dem bei gleichem Druck die höhere Temperatur um mindestens 50°C über der Kontaktierungstemperatur, vorzugsweise zwischen 80 und 120°C über der Kontaktierungstemperatur liegt, oder bei dem, bei gleicher Temperatur, der niedrigere Druck mindestens 5,6 mal niedriger, vorzugsweise zwischen 8 und 16 mal niedriger ist als der Kontaktierungsdruck.


Verfahren nach einem der vorhergehenden Ansprüche, bei dem das Gas aus der Vergasung von Biomasse, organischen Abfällen, Kohle oder einer Kombination davon stammt.

Revendications

1. Procédé de clarification d’un flux de gaz contenant des composants analogues à du goudron ou constitutifs du goudron comprenant des composés aromatiques ayant 6 atomes de carbone ou plus, comprenant la mise en contact du gaz avec un aryl polysiloxane organique liquide comme liquide de lavage.

2. Procédé selon la revendication 1, dans lequel le polysiloxane comporte une moyenne de 0,2 à 1,8 groupe aryle en C5-C14 par atome de silicium. 10

3. Procédé selon la revendication 1 ou 2, dans lequel le polysiloxane comporte une moyenne de 0,5 à 1,5 groupe aryle en C5-C10 et de 0,5 à 1,5 groupe alkyle en C1-C4 par atome de silicium.

4. Procédé selon l’une quelconque des revendications 1 à 3, dans lequel le polysiloxane a une masse molaire comprise entre 700 et 7 000 Da.

5. Procédé selon l’une quelconque des revendications 1 à 3, dans lequel le polysiloxane est un polyméthyléthylsiloxane ou un poly-diphényl-diméthyl-20 siloxane.


7. Procédé selon l’une quelconque des revendications précédentes, dans lequel le gaz doit être clarifié à partir des composants analogues à du goudron comprenant des composés aromatiques polycycliques ayant 9 à 18 atomes de carbone.

8. Procédé selon l’une quelconque des revendications précédentes, dans lequel le gaz est mis en contact avec le polysiloxane à une température de mise en contact comprise entre 30 et 150°C, de préférence entre 60 et 120°C, à une pression de mise en contact atmosphérique, ou à une température et une pression supérieures correspondantes.

9. Procédé selon la revendication 8, dans lequel le polysiloxane, après la mise en contact avec le gaz, est chauffé à une température qui est supérieure à la température de l’étape de mise en contact et/ou est dépressurisé à une pression qui est inférieure à la pression de mise en contact, de préférence en présence d’un gaz de stripage, et est ensuite renvoyé pour un cycle de contact 15 supplémentaire.

10. Procédé selon la revendication 9, dans lequel, à une pression égale, la température supérieure est supérieure d’au moins 50°C à la température de mise en contact, de préférence entre 80 et 120°C supérieure à la température de mise en contact, ou dans lequel, à température égale, la pression inférieure est au moins 5,6 fois, de préférence entre 8 et 16 fois inférieure à la pression de mise en contact.

11. Procédé selon l’une quelconque des revendications précédentes, dans lequel le gaz, avant mise en contact avec le polysiloxane, est traité avec un liquide de lavage comprenant des hydrocarbures aromatiques, à une température comprise entre 150 et 900°C.

12. Procédé selon l’une quelconque des revendications précédentes, dans lequel le gaz, avant mise en contact avec le polysiloxane, est soumis à un filtre électrostatique et / ou un fixateur aérosol.

13. Procédé selon l’une quelconque des revendications précédentes, dans lequel le gaz provient de la gazéification de la biomasse, de déchets organiques, de charbon ou d’une combinaison de ceux-ci.
REFERENCES CITED IN THE DESCRIPTION

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