

STAGED COMPLEMENTARY PSA SYSTEM FOR LOW ENERGY FRACTIONATION OF MIXED FLUID

FIELD

[0001] The present invention relates to low energy fractionation of a mixed fluid by using a staged complementary pressure swing adsorption system.

BACKGROUND

[0002] There is a growing need for the creation of energy efficient processes for the separation of multicomponent mixtures. The need for energy efficiency in this context has been augmented recently by environmental concerns. The separation of multicomponent mixtures is desirable for many reasons, most notably for use and production of a variety of industrial products. The distillation of propane and propylene illustrates this problem. This process is well known to be a very energy-intensive process and poses serious environmental risk.

[0003] Recent advances in material sciences have brought promise to this area, as a plethora of new materials have become available that can be used to selectively adsorb certain molecules from a gas. Modern adsorption processes have taken advantage of these materials as many of these processes employ the use of two different adsorption materials to separate multicomponent mixtures in the air.

[0004] Pressure swing adsorption (PSA) relies on swinging or cycling pressure over a bed of adsorbent through a range of values. In PSA processes, a gaseous mixture is conducted under pressure for a period of time over a first bed of a solid sorbent that is selective, or relatively selective, for one or more components, usually regarded as a contaminant, to be removed from the gaseous mixture. For example, a feed can be introduced into a PSA apparatus at a feed pressure. At the feed pressure, one or more of the components (gases) in the feed can be selectively (or relatively selectively) adsorbed, while one or more other components (gases) can pass through with lower or minimal adsorption. A component (gas) that is selectively adsorbed can be referred to as a "heavy" component of a feed, while a gas that is not selectively adsorbed can be referred to as a "light" component of a feed. For convenience, a reference to the "heavy" component of the feed can refer to all components (gases) that are selectively adsorbed, unless otherwise specified. Similarly, a reference to the "light" component can refer to all components (gases) that are not selectively adsorbed, unless otherwise specified. After a period of time, the feed flow into the PSA apparatus can be stopped. The feed flow can be stopped based on a predetermined schedule, based on detection of breakthrough of one or more heavy components, based on adsorption of the heavy component(s) corresponding to at least a threshold percentage of the total capacity of the adsorbent, or based on any other convenient criteria. The pressure in

the reactor can then be reduced to a desorption pressure that can allow the selectively adsorbed component(s) (gas(es)) to be released from the adsorbent. Optionally, one or more purge gases can be used prior to, during, and/or after the reduction in pressure to facilitate release of the selectively adsorbed component(s) (gas(es)). Depending on its nature, a full PSA cycle can optionally be performed at a roughly constant temperature. As PSA is usually enabled by at least adsorption and usually occurs on gaseous components, the terms "adsorption"/" adsorbent" and "gas(es)" are used as descriptors in the instant specification and claims, without intending to be limiting in scope, even though "absorption"/"absorbent"/"sorbent"/"sorption" and "component(s)" may be more generally applicable.

[0005] Multiple beds can be used to enable a complete cycle, where typically every bed sequentially goes through the same cycle. When a first PSA reactor satisfies a condition, such as the adsorbent in the reactor becoming sufficiently saturated, the feed flow can be switched to a second reactor. The first PSA reactor can then be regenerated by having the adsorbed gases released. To allow for a continuous feed flow, a sufficient number of PSA reactors and/or adsorbent beds can be used so that the first PSA reactor is finished regenerating prior to at least one other PSA reactor satisfying the condition for switching reactors.

[0006] U.S. Patent No. 4,744,803 discusses the use of a four-step PSA cycle. The adsorption bed is first pressurized by entry of gas from the bottom of the bed while the top end of the bed is closed. This is referred to as the pressurization step. The next step is high pressure feed, wherein feed gas enters under pressure from the top of the column and effluent is allowed to escape from the bottom of the column. At the conclusion of this step the column is closed at both ends and the pressurized gas is then released by opening the top end of the column. This is referred to as the blow down step. After the pressure has been reduced to a predetermined level by blowdown, the column is next purged of remaining product by feeding recycled product gas into the bottom end of the bed and allowing the gas remaining in the column to be forced out of the top end as effluent. This step would normally be terminated at the point where the purging gas reaches the top end of the column. The effluents from the blowdown and purge steps contain the component adsorbed by the column. This is generally referred to as the secondary product of the column. The primary product is the component or component which pass through the bed unadsorbed, i.e., the high pressure feed effluent.

[0007] U.S. Patent No. 4,744,803 also provides a PSA system where adsorbents 1 and 2 are disposed in 4 beds, which are respectively selective for species A and B. This system uses the secondary product of one bed (the blow down and purge effluent) partially or wholly as the feed for another bed containing a different, or complementary adsorbent. This system, however, has

the disadvantage that compressors are required to feed the secondary product of one feed to another. Thus greatly increasing the energy required to operate the system.

[0008] Another process known in the art is dual reflux PSA, described by Diagne et al., Ind. Eng. Chem. Res., Vol. 34 No. 9, 1995. In this process, an intermediate feed position divides an adsorber column into rectifying and stripping sections, and this process relies on external compression to create a constant product.

SUMMARY

[0009] The references cited above describe the use of compressors in PSA adsorption systems, and the use of compressors increases energy costs. The reduction or elimination of compressors from a PSA adsorption system would result in a significant decrease in energy costs.

[0010] Certain embodiments of the present invention are directed toward a pressure swing adsorption swing system for the fractionation of a multicomponent fluid comprising a first adsorbent column selective for a first component, a second adsorbent column selective for the first component, a third adsorbent column selective for a second component, a fourth adsorbent column selective for the second component; wherein said columns are fed with the multicomponent fluid while pressurized at a high pressure during at least one phase of a pressure swing adsorption cycle that enables a continuous supply of both a highly concentrated first component and a highly a concentrated second component to be recovered throughout the pressure swing adsorption cycle; wherein said columns are depressurized from the high pressure to a first intermediate pressure during at least one phase of the pressure swing adsorption cycle; wherein said columns are purged while pressurized at a second intermediate pressure during at least one phase of the pressure swing adsorption cycle and an intermittent supply of first and second component is created; wherein said columns are depressurized from a third intermediate pressure to a low pressure during at least one phase of the pressure swing adsorption cycle; wherein said columns are purged while pressurized at the low pressure during at least one phase of the pressure swing adsorption cycle by using an effluent from a column selective for a different component or from said intermittent supply; wherein said columns are re-pressurized to the high pressure during at least one phase of the pressure swing adsorption cycle.

[0011] Additional embodiments are directed toward a method for fractionation of a multicomponent stream having first and second components and for providing a continuous purified supply of said multicomponent stream using a pressure swing adsorption system having first adsorption columns selective for a first component and second adsorption columns selective for a second component, which comprises of steps of: feeding an adsorption column, selective for the first component, at a high pressure with a stream of the multicomponent gas and generating a

product comprising the second component; depressurizing said column to a first intermediate pressure, which is between the high pressure and a low pressure; purging said column at a second intermediate pressure by using one of the product of the second component or from the purge effluent of a column selective for the second component; depressurizing said column from a third intermediate pressure to the low pressure; purging said column at the low pressure by using product of second type of component; and increasing the pressure of said column from to the high pressure.

[0012] Further embodiments of the present invention are directed toward a method for fractionation of a multicomponent stream having first and second components and for providing a continuous purified supply of said multicomponent stream using a thermal swing adsorption system, wherein first adsorption columns are used that are selective for the first component and second adsorption columns are used that are selective for the second component, which comprises of steps of: feeding an adsorption column, selective for the first component, at a first high temperature with a stream of the multicomponent stream and generating a product comprising the second component; decreasing the temperature of said column by coming to thermal equilibrium with another column which is at a first low temperature; purging said column at a first intermediate temperature by using one of the product comprising the second component or the purge effluent of a column selective for the second component; purging said column at a second low temperature by using the product comprising the second component; and increasing the temperature of said column from a third low temperature to a third high temperature, by coming to thermal equilibrium with another column which is at higher temperature than said column.

[0013] The invention may be embodied by numerous other devices and methods. The description provided herein, when taken in conjunction with the annexed drawings, discloses examples of the invention. Other embodiments, which incorporate some or all steps as taught herein, are also possible.

BRIEF DESCRIPTION OF THE DRAWINGS

[0014] FIG. 1 shows a flow chart depicting the various stages in one cycle for an exemplary four column, staged complementary PSA system for separating components A and B from a feed at minimum energy cost.

[0015] FIG. 2 shows an exemplary adsorbent bed that permits heat transfer between two adsorbent columns.

[0016] FIG 3 shows an exemplary compact adsorbent bed that permits heat transfer between four adsorbent columns.

[0017] FIG. 4 shows a flow chart depicting the various stages in another exemplary cycle for a four column, staged complementary PSA system used to produce a continuous highly concentrated CO₂ and N₂ product at minimum energy cost.

[0018] FIG. 5 shows a flow chart depicting the various stages in another exemplary cycle for a four column, staged complementary PSA system used to produce a continuous highly concentrated CH₄ product from a mixture of N₂ and CH₄ at minimum energy cost.

DETAILED DESCRIPTION

[0019] In various aspects, systems and methods are provided for low energy fractionation of a mixed fluid by using a staged complementary pressure swing adsorption system. In these embodiments intermediate staged pressure steps are provided in a multi-bed PSA system to enable the pressurization and purging of complementary adsorbent beds such that no compressors are required.

[0020] The first embodiment will be described with reference to FIG. 1. The embodiment includes a four bed PSA system, where two of the beds are selective or relatively selective for one of the components of a mixture, while the other two beds are selective or relatively selective for the other component of the mixture. The adsorbent columns (1+) and (2+) are selective for component A, and the adsorbent columns (1-) and (2-) are selective for component B. As will be described, a continuous supply of component A and B product is generated throughout the cycle.

[0021] It should be appreciated that the exact timings and pressure of the cycle will vary depending on the particulars of the process and materials used, e.g., the component that is adsorbed and the adsorbent being used. Some examples of molecule combinations this system can be used to separate include nitrogen/methane, nitrogen/carbon dioxide, hydrogen/carbon dioxide, methane/carbon dioxide, argon/oxygen, oxygen/nitrogen, propane/propylene, ethane/ethylene, etc. Depending on the molecules to be separated, different adsorbent materials are used. Such adsorbent materials may include zeolites such as ITQ/H-mordenite for the separation of nitrogen/methane, for example. Other complementary pairs of adsorbents are well known in the art.

[0022] Examples of suitable adsorbent materials selective will now be described. Such material includes a mixed metal oxide adsorbent, such as an adsorbent including a mixture of an alkali metal carbonate and an alkaline earth metal oxide and/or a transition metal oxide. Examples of suitable alkali metal carbonates can include, but are not limited to, a carbonate of lithium, sodium, potassium, rubidium, cesium, or a combination thereof, e.g., a carbonate of lithium, sodium, potassium, or a combination thereof. Examples of suitable alkaline earth metal

oxides can include, but are not limited to, oxides of magnesium, calcium, strontium, barium, or a combination thereof, *e.g.*, oxides of magnesium and/or calcium. Some examples of suitable transition metal oxides can include, but are not limited to, oxides of lanthanide series metals, such as lanthanum, and/or of transition metals that can form oxides with the metal in a +2 or +3 oxidation state (such as yttrium, iron, zinc, nickel, vanadium, zirconium, cobalt, or a combination thereof).

[0023] In some aspects, the carbonate can be selected independently from the oxide in the mixed metal oxide. In such aspects, the carbonate can include, consist essentially of, or be lithium carbonate, sodium carbonate, potassium carbonate, rubidium carbonate, and/or cesium carbonate (*e.g.*, lithium carbonate, sodium carbonate, and/or potassium carbonate; lithium carbonate and/or potassium carbonate; lithium carbonate and/or sodium carbonate; or sodium carbonate and/or potassium carbonate).

[0024] In aspects where the carbonate is selected independently from the oxide, the oxide can be an alkaline earth oxide, a transition metal oxide, a combination of two or more alkaline earth oxides, a combination of two or more transition metal oxides, or a combination of oxides including at least one alkaline earth oxide and at least one transition metal oxide. In aspects where the independently selected oxide includes one or more alkaline earth oxides, a suitable alkaline earth oxide can include, consist essentially of, or be magnesium oxide, calcium oxide, strontium oxide, and/or barium oxide, *e.g.*, including at least magnesium oxide and/or calcium oxide.

[0025] In aspects where the independently selected oxide includes one or more transition metal oxides, suitable transition metals can include, consist essentially of, or be one or more transition metals that can form oxides with the metal in a +2 or +3 oxidation state (*e.g.*, yttrium oxide, iron oxide, zinc oxide, nickel oxide, vanadium oxide, cobalt oxide, zirconium oxide, lanthanum oxide, other oxides of lanthanide metals, and/or a combination thereof). One preferred option includes a transition metal oxide selected from lanthanum oxide and/or zirconium oxide. Another option includes a metal oxide selected from lanthanum oxide, yttrium oxide, zirconium oxide, and/or zinc oxide. Yet another option includes a metal oxide selected from nickel oxide, cobalt oxide, and/or iron oxide. Mixtures within each of these options and/or across options are also contemplated, such as mixtures of lanthanum oxide with zinc oxide and/or vanadium oxide; mixtures of lanthanum oxide with iron oxide, cobalt oxide, and/or nickel oxide; mixtures of zirconium oxide with yttrium oxide, zinc oxide, and/or vanadium oxide; and mixtures of zirconium oxide with iron oxide, cobalt oxide, and/or nickel oxide.

[0026] In aspects where the independently selected oxide includes one or more alkali metal oxides and one or more transition metal oxides, suitable alkali metal oxides can include, consist essentially of, or be magnesium oxide, calcium oxide, strontium oxide, and/or barium oxide, while suitable transition metals can include, consist essentially of, or be transition metals that can form oxides with the metal in a +2 or +3 oxidation state, such as yttrium oxide, iron oxide, zinc oxide, nickel oxide, vanadium oxide, cobalt oxide, zirconium oxide, lanthanum oxide, and/or other lanthanide oxides. Each of these alkali metal oxides and transition metal oxides can be independently selected individually or in any combination of multiple transition metal oxides. Examples of mixtures can include, consist essentially of, or be a mixture of oxides where at least one oxide is lanthanum oxide, zirconium oxide, and/or magnesium oxide; a mixture of oxides where the mixture includes at least two of lanthanum oxide, zirconium oxide, and magnesium oxide; a mixture of oxides where one oxide is magnesium oxide and/or calcium oxide; and/or a mixture of oxides where at least one oxide is lanthanum oxide, yttrium oxide, and/or zirconium oxide.

[0027] In some alternative aspects, a mixed metal oxide can include an alkaline earth carbonate in combination with a transition metal oxide. In such aspects, the alkaline earth carbonate can include, consist essentially of, or be magnesium carbonate and/or calcium carbonate. Additionally or alternately, the alkaline earth carbonate can be present in a mixture with an alkali metal carbonate. Examples of such carbonate mixtures can include, consist essentially of, or be mixtures of lithium carbonate with magnesium carbonate, lithium carbonate with calcium carbonate, potassium carbonate with magnesium carbonate, potassium carbonate with calcium carbonate, sodium carbonate with magnesium carbonate, and sodium carbonate with calcium carbonate (*e.g.*, lithium carbonate with magnesium carbonate or potassium carbonate with magnesium carbonate). In such aspects, suitable transition metals can include, consist essentially of, or be transition metals that can form oxides with the metal in a +2 or +3 oxidation state, such as yttrium oxide, iron oxide, zinc oxide, nickel oxide, vanadium oxide, cobalt oxide, zirconium oxide, lanthanum oxide, other lanthanide oxides, and/or a combination thereof. Each of these alkaline earth carbonates and transition metal oxides can be independently selected individually or in any combination of multiple alkaline earth carbonates and/or multiple transition metal oxides. For the transition metal oxide, one preferred option can include a transition metal oxide selected from lanthanum oxide or zirconium oxide. Another option can include a metal oxide selected from lanthanum oxide, yttrium oxide, zirconium oxide, and/or zinc oxide. Yet another option can include a metal oxide selected from nickel oxide, cobalt oxide, and/or iron oxide. Mixtures within each of these options and/or across options are also

contemplated, such as mixtures of oxides where at least one oxide is lanthanum oxide and/or zirconium oxide; mixtures of lanthanum oxide with zinc oxide and/or vanadium oxide; mixtures of lanthanum oxide with iron oxide, cobalt oxide, and/or nickel oxide; mixtures of zirconium oxide with yttrium oxide, zinc oxide, and/or vanadium oxide; and/or mixtures of zirconium oxide with iron oxide, cobalt oxide, and/or nickel oxide.

[0028] Additional or alternative materials can include hydrotalcites.

[0029] Additionally or alternatively, the present invention can include or more of the following embodiments.

[0030] A PSA cycle of column (1+) will now be described. The cycle of column (1+) commences with a high pressure ("PHIGH") feed step, during which feed gas having at least two components A and B is supplied to a feed end of the PSA reactor vessel at PHIGH. An example of a feed gas is the recycled turbine exhaust gas from a low emission power generation system. As the feed gas is supplied to a feed line, the "product," i.e., component B, is generated from a product side, as the component A is adsorbed into the (1+) bed.

[0031] After the high pressure feed step, column (1+) is be depressurized to middle pressure ("PMID"). PMID is an intermediate pressure, which is less than PHIGH, but more than a low pressure ("PLOW"). Depressurization enables the adsorbed gas molecules to be released from the adsorbent material in the column. As the pressure of a column decreases, the amount of released adsorbent gas increases in the column. The released adsorbent gas from a column will be referred to as effluent gas. Next, the released effluent gas of column (1+) is purged with the high pressure feed of column (2+). At the same time, column (2-) (also at intermediate pressure) is purged with this effluent, received from column (1+). The communication between columns (1+) and (2-) creates an intermediate A and B product. This intermittent product may be used as product or stored in appropriate storage tanks, and may be used elsewhere in the cycle as purge gas during low-pressure purges. It will be appreciated that the cycle is advantageously performed without the use of compressors.

[0032] Next, column (1+) is depressurized to PLOW. After depressurization, column (1+) is purged using the effluents of the complementary column (2-). Purging facilitates the release of the adsorbed component A in column (1+). In this way, no external compression is needed to provide the purge gas for column (1+). Alternatively, column (1+) may be purged by using the effluent of a complementary column (1-) or (2-) stored in an appropriate storage tank at PMID. After purging, column (1+) is re-pressurized to PHIGH. Once re-pressurization is complete, feed gas fed into column (1+) at high pressure during a high feed pressure step. This enables product B to be generated, as component A from the feed gas is adsorbed into column (1+).

[0033] The re-pressurization marks the end of the first half of the cycle for column (1+). The second half of the cycle of column (1+) is sequenced in generally the same way as the first half of the cycle, with two exceptions. First, after column (1+) is depressurized to PMID, the column (2-) is purged with the effluent of column (1+) to make an intermittent A and B product rather than column (1-). Second, column (1+) is purged at low pressure by using the purge effluent from column (1-) rather than column (2-).

[0034] Next a cycle of column (2+) will be described in reference to the timing of phase 1. Column (2+), like column (1+), is an adsorbent bed selective for component A. As column (1+) commences a high pressure feed step in the cycle, column (2+) is depressurized from PHIGH to PMID. After depressurization, column (2+) is purged with the high pressure feed. The effluent of column (2+) is fed to column (1-), and an intermittent A and B product is created. The product may be stored in an appropriate storage tank and may be used later in the cycle to purge a column at low pressure. The column (2+) is next depressurized from PMID to PLOW, and the remaining adsorbed gas in column (2+) may be purged with the effluent gas of column (2-). This enables column (2+) to be purged without the need for external compressors.

[0035] After purging column (2+) of the remaining adsorbed gas, the pressure of column (2+) is re-pressurized to PHIGH. Column (2+) is then fed with feed gas, and the product, i.e., component B, is generated as component A from the feed gas is adsorbed into the column (2+). After this high pressure feed step in column (2+), the pressure of the column is decreased to PMID.

[0036] This step concludes the first half of the cycle of column (2+). The second half of the cycle of column (2+) is sequenced in the same way as the first half of the cycle, with two exceptions. First, when column (2+) is depressurized to PMID, column (2-) rather than (1-), is purged with the effluents of column (1+), to make an intermittent A and B. Second, when column (2+) is purged at low pressure, it is fed with the purge effluent of column (1-) rather than column (2-). It will be appreciated that column (1+) continuously generates component B product until column (2+) commences this high pressure feed step. This enables product B to be generated at all times during this cycle. Product B will be generated from either column (1+) or (2+) throughout the cycle.

[0037] The cycle of adsorbent column (1-) will be described. In FIG. 1, column (1-) commences the cycle at a re-pressurization step, from PLOW to PMID. After this step, column (1-) is fed the purge effluent of column (2+), which is at PMID, to create an intermittent product A and B. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks.

[0038] Next, column (1-) is re-pressurized to PHIGH. After this re-pressurization, feed gas is supplied to the feed end of the column (1-) at PHIGH. In this step, product A can be generated as component B is adsorbed in the column from the feed gas. Following this high feed pressure step, the pressure of column (1-) is reduced to PMID. Afterward, column (1-) is fed with the purge effluent of column (1+), also at PMID, to create an intermittent product A and B. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks.

[0039] Next, column (1-) is re-pressurized to PHIGH. After re-pressurization, feed gas is supplied to the feed end of the column (1-) at PHIGH. In this step product A can be generated as component B is adsorbed in the column. This high pressure feed step is repeated, such that the second high pressure feed step of column (1-) begins as the second high pressure feed step of column (1+) begins. Column (1-) is then depressurized to PLOW. After depressurization, Column (1-) is purged with high pressure product of column (2-), or alternatively, using the intermittent product stored at PMID. The effluent from column (1-) created by this purge is fed to purge column (2+) at low pressure.

[0040] After column (1-) is purged, the column is re-pressurized to PHIGH. Next, feed gas is supplied to the feed end of column (1-) at PHIGH. In this step component A can be generated as component B is adsorbed in the column. Column (1-) is then depressurized to PLOW. Column (1-) is then depressurized to PLOW. After depressurization, Column (1-) is purged with high pressure product of column (2-), or alternatively, using the intermittent product stored at PMID. The effluent from column (1-) created by this purge is fed to purge column (1+) at low pressure.

[0041] The cycle of column (2-) will now be described. FIG. 1 commences with a high pressure feed step, where product A is generated and component B is adsorbed by column (2-). Next, column (2-) is depressurized to PLOW. After depressurization, Column (2-) is purged using the high pressure product of column (1-), or alternatively, using the intermittent product stored at PMID. The effluent from column (2-) created by this purge is fed to purge column (2+) at low pressure.

[0042] After, column (2-) is re-pressurized to PHIGH. Following this step, a high-pressure feed step occurs, where product A is generated and component B is adsorbed by column (2-). Column (2-) is then depressurized to PLOW. After depressurization, Column (2-) is purged using the high pressure product of column (1-), or alternatively, using the intermittent product stored at PMID. The effluent from column (2-) created by this purge is fed to purge column (1+) at low pressure.. Column (2-) is then re-pressurized to PMID. After this step, column (2-) is fed

the purge effluent of column (2+), which is at PMID, to create an intermittent product A and B. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks.

[0043] Column (2-) is then re-pressurized to PHIGH. Following this step, a high-pressure feed step occurs, where product A is generated and component B is adsorbed by column (2-). After this, column (2-) is de-pressurized to PMID. Next, column (2-) is fed the purge effluent of column (1+), which is at PMID, to create an intermittent product A and B. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks.

[0044] After this, column (2-) is re-pressurized to PHIGH. This is followed by a high-pressure feed step, where product A is generated, i.e. component A, and component B is adsorbed by column (2-). It should be appreciated that at any time column (2-) is generating product A during a high pressure feed step, column (1-) is not. Likewise, whenever column (1-) is generating product A in a high-pressure feed step, column (2-) is not. This cycle has the advantage of generating a continuous supply of product A.

[0045] It is appreciated that the pressurization and re-pressurization steps in the first embodiment may be accomplished through "equalizations of pressures" between two different complementary columns. Connecting the columns together, opening the valves on the connecting ends of the columns, and then closing the valves on the other side of the columns enable the equalization of pressure between two complementary columns (i.e. columns of different adsorbent materials). Equalizations of pressure in this matter can increase the energy efficiency of the cycle.

[0046] It will be appreciated that the pressures PHIGH, PMID, and PLOW described herein are relative pressures, and the value of each may be adjusted based on the particular components and cycles at hand.

[0047] In a second embodiment of this invention, the re-pressurization and depressurization steps discussed in FIG. 1 are replaced with cooling and heating steps, respectively. During these cooling and heating steps, the pressure of the columns is held nominally the same. The heating and cooling steps have the same effect (as pressure is kept constant) as the re-pressurization and depressurization steps in the first embodiment.

[0048] FIG. 2 shows an example of how the complementary adsorbents can be deployed in an adsorbent column that permits heat transfer between adsorbents. In FIG. 2, two adsorbent beds are placed in a partitioned vessel with common walls. The adsorbent beds are placed

coaxially in pairs. Such a divided wall adsorption column offers the advantage of compactness while enabling the heat transfer between adsorbents.

[0049] FIG 3 shows an exemplary compact adsorbent bed that may be used in this invention, which permits heat transfer between four adsorbent columns. FIG. 3 depicts a configuration of the adsorbent columns where they are placed in a partitioned vessel with common walls. The adsorbent columns are placed coaxially in quadruples. The compactness of the adsorbent bed in Figure 3 additionally allows efficient pressure equalizations between different adsorbent columns (e.g. 1+, 2 -) as the columns are situated in close proximity to one another other and thus do not require the use of expensive piping. For the same reason, the effluent of one column can be efficiently discharged to another. The partition separating the adsorbent columns in Figure 3 may be made of a thermal conducting material such that heat transfer is permitted between adjacent columns.

FIRST EXAMPLE

[0050] A first illustrative example of the deployment of the novel staged complementary PSA cycle in FIG. 3 is now discussed for the separation of a power generation stream of CO₂/N₂. Two vessels (denoted as columns 1+ and 2+, respectively) of unit cross-sectional area (1 meter squared) and length of 2 meters are filled with an adsorbent selective for N₂ as compared to CO₂. A further two vessels (denoted as columns 1- and 2-, respectively) also of unit cross-sectional area (1 meter squared) and a length of 2 meters are filled with an adsorbent selective for CO₂ as compared to N₂.

[0051] The adsorbent material of columns (1+) and (2+) is selective for N₂ and may be composed of 4A, NaA zeolite or of LiX zeolite. The adsorbent material of columns (1-) and (2-) is selective for carbon dioxide and may be composed of activated carbons, aluminas, metal organic framework (MOF), mixed metal oxides, hydrotalcites, or a combination of these. The process described in reference to FIG. 3 uses no additional compression and produces an enriched carbon dioxide product containing 93.8 mol % CO₂ at a recovery of 70% and an N₂ enriched product containing 97.7 mol % N₂ at a recovery of 98.8%.

[0052] In accordance with the cycle shown in FIG. 3, columns (1+) and (2+) are subjected to a feed step. The feed is composed of 10% CO₂ and 90% N₂, and is provided at a pressure of 25 bar and temperature of 250 Celsius at a feed rate of 630 kg/hr. Columns (1-) and (2-) are subjected to a feed step with a composition of 10% carbon dioxide and 90% N₂ at a pressure of 25 bar and a temperature of 250 °C at a feed rate of 350 kg/hr. Various pressures and times associated with the cycle steps are depicted in FIG. 3 and will be described hereafter.

[0053] The cycle of column (1+) will now be described. Column (1+) is first pressurized at 25 bar and fed with a gaseous mixture of carbon dioxide and nitrogen for 260 seconds. N₂ is adsorbed in this column, while air with a high concentration of CO₂ is generated from the column. Next, column (1+) is depressurized for 30 seconds and brought to a pressure of 23 bar. Simultaneously, column (1+) is connected to column (1-) such that the pressure between the two columns is equalized. The pressure of column (2-) is brought from a pressure of 12 bar to 23 bar. After, the pressure of column (1+) is held for 15 seconds at 22.7 bar. This can be accomplished by closing all of the valves in column (1+). Next, the pressure between column (1+) is equalized with column (1-) for 15 seconds, raising the pressure back up to 23 bar.

[0054] Following this step, column (1+) is depressurized to 17.4 bar and then purged using the high pressure product from column (2+). It is appreciated that the effluent from column (1+) in this step is fed to purge column (1-), which create an intermittent CO₂ and N₂ product at an intermediate pressure (17.4 bar). This illustrates the way that the present invention can purge product without using a compressor.

[0055] Column (1+) is then brought to a low pressure, of 1.3 bar, and this step takes approximately 15 seconds. Following this, column (1+) may be purged at this low pressure 45 seconds from the high pressure product of column (2+), alternatively purging could occur from stored intermittent product. Column (1+) next is brought to a pressure of 10 bar, for 5 seconds. This is accomplished by feeding column (1+) with the stream from columns (1-) and (2-) at high pressure. After this, the pressure is raised to 12 bar by equalizing the pressure of column (1+) with column (2-) for 12 seconds. Thus, the pressure of column (1+) is raised from 1.3 bar to 12 bar without using a compressor.

[0056] Column (1+) is next re-pressurized to 25 bar by connecting the column with the feed gas for 25 seconds. Afterwards, column (1+) generates highly concentrated carbon dioxide for 260 seconds during a high feed pressure step at 25 bar of pressure. Next, column (1+) is depressurized for 30 seconds and brought to a pressure of 23 bar. The pressure is held constant in column (1+) at 22.7 bar for the next 15 seconds

[0057] Simultaneously, column (1+) is connecting to column (1-) such that the pressure of column (1-) can be increased. During this time period the pressure of column (1-) is brought from a pressure of 12 bar to 23 bar. After this, the pressure of column (1+) is held for 15 seconds at 23 bar. This can be accomplished by closing all of the valves of column (1+). Following this step, column (1+) is connected to column (2-). For 15 seconds, the pressure between the columns is equalized to an intermediate pressure, around 23 bar.

[0058] Following this step, column (1+) is depressurized to 17.4 bar and then purged using the high pressure product from column (2+). It is appreciated that the effluent from column (1+) in this step is fed to purge column (2-), which create an intermittent CO₂ and N₂ product at an intermediate pressure (17.4 bar). Column (1+) is then depressurized to a low pressure of 1.3 bar. This step takes approximately 15 seconds. Following this, column (1+) is purged for 45 seconds using the high pressure product of column (2+), or, alternatively, with stored intermittent product. Column (1+) can be purged without the use of a compressor. Column (1+) is then brought to a pressure of 10 bar for 5 seconds. This is accomplished by feeding column (1+) with the stream from columns (1-) and (2-) at high pressure. After this, the pressure is raised to 12 bar by equalizing the pressure of column (1+) with column (2-) for 15 seconds. Thus, the pressure of column (1+) is raised from 1.3 bar to 12 bar without the use of compressors. Finally, column (1+) is re-pressurized to 25 bar.

[0059] The cycle of column (2+) will now be described. The cycle of column (2+) commences as column (2+) is depressurized from 25 bar to 23 bar over a 30 second period. Simultaneously, column (2+) is connected to column (1-), such that the pressure of column (1-) can be increased while the pressure of column (2-) decreases. During this time period the pressure of column (2-) is brought from a pressure of 12 bar to 23 bar. After this step, the pressure of column (2+) is held for 30 seconds at 23 bar. This can be accomplished by closing all of the valves of column (2+). Following this step, column (2+) is purged using the high pressure product from column (1+). It is appreciated that the effluent from column (2+) in this step is fed to purge column (1-), which creates an intermittent CO₂ and N₂ product at an intermediate pressure. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks. During this step the pressure is decreased to 10 bar.

[0060] The pressure of column (2+) is then further decreased to 1.3 bar. This step takes approximately 15 seconds. Next, column (2+) is purged at this low pressure for 45 seconds using the high pressure product of column (1+). After, column (2+) can be brought to a pressure of 10 bar for 5 seconds. This is accomplished by feeding column (2+) with the stream from columns (1-) and (2-) at high pressure and closing the other valve of column (2+). The pressure of column (2+) is further raised to 12 bar by equalizing the pressure of column (2+) with column (2-) for 15 seconds. Thus, column (2+) is purged and the pressure of column (2+) is raised from 1.3 bar to 12 bar without the use of compressors.

[0061] Column (2+) is next re-pressurized to 25 bar. Column (2+) then generates highly concentrated carbon dioxide for 260 seconds during the high feed pressure step at 25 bar of

pressure. Next, column (2+) is depressurized for 30 seconds and brought to a pressure of 23 bar. Simultaneously, column (2+) is connected to column (2-) such that the pressure of the two is equalized. During this time period the pressure of column (2-) is brought from a pressure of 12 bar to 23 bar. After this equalization step, the pressure of column (2+) is held for 15 seconds at 23 bar. This can be accomplished by closing all of the valves on column (2+).

[0062] Following this step, column (2+) is purged using the high pressure product from column (1+). It is appreciated that the effluent from column (2+) in this step is fed to purge column (2-), which creates an intermittent CO₂ and N₂ product at an intermediate pressure. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks. During this step the pressure is decreased to 10 bar.

[0063] Column (2+) is then depressurized to a low pressure of 1.3 bar, which takes approximately 15 seconds. Next, column (2+) is purged for 45 seconds using the high pressure feed of column (1+), or, alternatively, the stored intermittent product. Column (2+) is then brought to a pressure of 10 bar for 5 seconds. This is accomplished by feeding column (2+) with the stream from columns (1-) and (2-) at high pressure. After this, the pressure is raised to 12 bar by equalizing the pressure of column (2+) with column (1-) for 15 seconds. Thus, purging and increasing the pressure of column is accomplished without the use of compressors.

[0064] Column (2+) is next re-pressurized to 25 bar. After this, column (2+) is held at 25 bar and fed with a gaseous mixture of CO₂ and N₂ for 260 seconds. N₂ is adsorbed in this column, while air with a high concentration of CO₂ can be generated from the column. As can be seen in FIG. 3 and just described, the cycle is coordinated in such a way that either columns (2+) and (1+) are in a high pressure step producing CO₂, thus CO₂n is continuously generated.

[0065] The cycle of column (1-) will now be described. Column (1-) commences the cycle configured to column (2+) such that the pressure of column (1-) equalizes with the pressure of column (2-). In this step, the pressure of column (1-) is increased from 12 bar to 23 bar after 30 seconds. Next, the pressure of column (1-) is held at a constant 23 bar pressure for 30 seconds.

[0066] Following this, column (1-) is fed with the purge effluent of (2+) (at 10 bar); this step create intermittent CO₂ and N₂ products at 10 bar. During this step, the pressure of column (1-) is decreased from 23 bar to 12 bar over the course of 60 seconds. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks.

[0067] Column (1-) is next re-pressurized from 12 bar to 25 bar over a period of 65 seconds. After this, column (1-) is held at a pressure of 25 bar and fed with the high-pressure feed of the gaseous mixture of CO₂ and N₂ for 120 seconds. CO₂ is adsorbed in this column and air with a

high concentration of N₂ is generated from the column. After this, column (1-) is depressurized to 23 bar over the course of 15 second. Column (1-) is then further depressurized to 17.4 bar and purged with the effluent of (2+) (at 17.4 bar); this step create intermittent CO₂ and N₂ products at 17.4 bar over the course of 60 seconds. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks.

[0068] Column (1-) is again re-pressurized from 12 bar to 25 bar over a period of 65 seconds. After this, column (1-) is held at a pressure of 25 bar and fed with the high-pressure feed of the gaseous mixture of CO₂ and N₂ for 260 seconds. CO₂ is adsorbed in this column and air with a high concentration of N₂ is generated from the column. Column (1-) is then connected to column (2+), such that the pressure between the columns is equalized to an intermediate pressure, 12 bar. This step takes 15 seconds. Next, the pressure of column (1-) is held at a constant 23 bar pressure for 60 seconds. This occurs until column (1-) is reconnected with column (1+) such that the pressure between the columns is equalized to an intermediate pressure, 23 bar for 30 seconds. Thus, the pressure of column (1-) is raised from 12 to 23 bar without the use of compressors.

[0069] Column (1-) is then re-pressurized from 23 bar to 25 bar over a period of 15 seconds. After this, column (1-) is held at a pressure of 25 bar and fed with the high-pressure feed of the gaseous mixture of CO₂ and N₂ for 140 seconds. Column (1-) is connected with column (1+) such that the pressure between the two can equalize to 12 bar. This takes 15 seconds. Finally, the pressure in column (1-) is held for 60 seconds.

[0070] The cycle of column (2-) will now be described. Column (2-) commences the cycle in a high pressure feed step where column (2-) is held at a pressure of 25 bar and fed with the high-pressure feed of the gaseous mixture of CO₂ and N₂ for 185 seconds. CO₂ is adsorbed in this column and air with a high concentration of N₂ is generated from the column. Column (2-) is then connected to column (2+), such that the pressure between the columns is equalized to an intermediate pressure, 12 bar. This takes 15 seconds. Next, the pressure of column (2-) is held at a constant 23 bar pressure for 60 seconds. This occurs until the column (2-) is reconfigured with column (1+) such that the pressure between the columns is equalized to an intermediate pressure, 23 bar for 30 seconds. Thus, the pressure of column (2-) is raised from 12 to 23 bar without the use of compressors.

[0071] Column (2-) is then re-pressurized from 23 bar to 25 bar over a period of 15 seconds. After this, column (2-) is held at a pressure of 25 bar and fed with the high-pressure feed of the gaseous mixture of CO₂ and N₂ for 140 seconds. CO₂ is adsorbed in this column and air with a high concentration of N₂ is generated from the column. Column (2-) is then connected to column

(1+), such that the pressure between the columns is equalized to an intermediate pressure, 12 bar. This takes 15 seconds. Next, the pressure of column (2-) is held at a constant 23 bar pressure for 60 seconds. This occurs until column (2-) is reconnected with column (2+), such that the pressure between the columns is equalized to an intermediate pressure of 23 bar for 30 seconds. This is briefly followed by a hold period, where the pressure remains unchanged in column (2-) for 23 seconds. Following this, column (2-) is purged with the effluent of (2+) (at 10 bar). This step creates intermittent CO₂ and N₂ products. During this step, the pressure of column (2-) is decreased from 23 bar to 10 bar over the course of 60 seconds.

[0072] Column (2-) is next re-pressurized from 10 bar to 25 bar over a period of 65 seconds. After this, column (2-) is held at a pressure of 25 bar and fed with the high-pressure feed of the gaseous mixture of CO₂ and N₂ for 120 seconds. CO₂ is adsorbed in this column and air with a high concentration of N₂ is generated from the column. Following this step, column (2-) is connected to columns (1-) and the pressure is equalized between them to 23 bar. This takes 15 seconds. Following this, column (2-) is depressurized to 17.4 bar and purged using the effluent of column (1-) at 17.4 bar. This takes 60 seconds and creates an intermittent CO₂ and N₂ product.

[0073] Column (2-) is then re-pressurized from 12 bar to 25 bar over a period of 65 seconds. After this, column (1-) is held at a pressure of 25 bar and fed with the high-pressure feed of the gaseous mixture of CO₂ and N₂ for 25 seconds. CO₂ is adsorbed in this column and air with a high concentration of N₂ is generated from the column. As can be seen in FIG. 3 and just described, columns (2-) and (1-) are configured in such a way that a high concentration of carbon dioxide is continuously produced throughout the cycle.

SECOND EXAMPLE

[0074] A second illustrative example of the deployment of the novel staged complementary PSA cycle in FIG. 4 is now discussed for the separation of CH₄/N₂. This process described hereafter may advantageously be used after power generation. Two vessels (denoted as columns 1+ and 2+, respectively) of unit cross-sectional area (1 meter squared) and length of 2 meters are filled with an adsorbent selective for CH₄ as compared to N₂. A further two vessels (denoted as columns 1- and 2-, respectively) also of unit cross-sectional area (1 meter squared) and a length of 2 meters are filled with an adsorbent selective for N₂ as compared to CH₄.

[0075] Adsorbent material more selective for CH₄ than N₂ can be chosen from H/Na Mordenite, activated carbons, aluminas, metal organic framework (MOF), mixed metal oxides, hydrotalcites or combinations thereof. Adsorbent materials more selective for N₂ than CH₄ can be chosen from molecular sieves such clinoptilolite, rho-zeolite or ITQ as well as carbon

molecular sieve. The columns (1+) and (2+) in this example are subjected to a feed step with the composition of 10% N₂ and 90% CH₄, a pressure of 30 bar, a temperature of 50° C and a feed rate of 486 kg/hr. The columns (1-) and (2-) in this example are subjected to a feed step with the composition of 10% N₂ and 90% CH₄, a pressure of 30 bar, a temperature of 50° C and a feed rate of 583 kg/hr. Various pressures and times associated with the cycle steps are as shown on in Figure 3

[0076] The second example uses no additional compression and produces a methane enriched product containing 95.8 mol % CH₄ at a recovery of 93.8 %. It will be appreciated that in order to produce this highly enriched constant product the ratio of feed time to total cycle time for a given adsorbent bed selective for CH₄ may be set between 0.3 to 0.7, and preferably is 0.5. The ratio for purge time to total cycle time for a given adsorbent bed selective for CH₄ is optimized between ranges of 0.15 and 0.25, and preferably set at 0.2.

[0077] The cycle may be configured to produce highly enriched CH₄ product by setting the ratio of feed times to total cycle times of a given column selective for N₂ to approximately between 0.6 and 0.4, and may be preferably set to 0.5. The ratio of purge times to total cycle times of a given column selective for N₂ is optimally set to approximately between 0.08 and 0.16, and may be preferably set to 0.12.

[0078] The cycle of column (1+) will now be described. Column (1+) is first pressurized at 40 bar and fed with a gaseous mixture of CH₄ and N₂ for 260 seconds. CH₄ is adsorbed in this column, while air with a high concentration of N₂ is generated from the column. Next, column (1+) is depressurized for 30 seconds and brought to a pressure of 34.2 bar. Simultaneously, column (1+) is connected to column (1-) such that the pressure between the two columns is equalized. The pressure of column (2-) is brought from a pressure of 20 bar to 34.2 bar. After, the pressure of column (1+) is held for 15 seconds at 34.2 bar. This can be accomplished by closing all of the valves in column (1+). Next, the pressure between column (1+) is equalized with column (1-) for 15 seconds, raising the pressure back up to 36 bar.

[0079] Following this step, column (1+) is depressurized to 24 bar and then purged using the high pressure product from column (2+). It is appreciated that the effluent from column (1+) in this step is fed to purge column (1-), which create an intermittent CH₄ and N₂ product at an intermediate pressure (24 bar). This illustrates the way that the present invention can purge product without using compressors.

[0080] Column (1+) is then brought to a low pressure, of 1.3 bar, this step takes approximately 15 seconds. Following this, column (1+) may be purged at this low pressure 45 seconds from the high pressure product of column (2+), alternatively purging could occur from

stored intermittent product. Column (1+) next is brought to a pressure of 13.8 bar, for 5 seconds. This is accomplished by feeding column (1+) with the stream from columns (1-) and (2-) at high pressure. After this, the pressure is raised to 20 bar by equalizing the pressure of column (1+) with column (2-) for 20 seconds. Thus, the pressure of column (1+) is raised from 1.3 bar to 20 bar without the use of compressors.

[0081] Column (1+) is next re-pressurized to 40 bar by connecting the column with the feed gas for 40 seconds. Afterwards, column (1+) generates highly concentrated N₂ for 260 seconds during a high feed pressure step at 40 bar of pressure. Next, column (1+) is depressurized for 30 seconds and brought to a pressure of 34.2 bar. The pressure is held constant in column (1+) at 34.2 bar for the next 15 seconds

[0082] Simultaneously, column (1+) is connecting to column (1-) such that the pressure of column (1-) can be increased. During this time period the pressure of column (1-) is brought from a pressure of 20 bar to 36 bar. After this, the pressure of column (1+) is held for 15 seconds at 34.2 bar. This can be accomplished by closing all of the valves of column (1+). Following this step, column (1+) is connected to column (2-). For 15 seconds, the pressure between the columns is equalized to an intermediate pressure, around 36 bar.

[0083] Following this step, column (1+) is depressurized to 24 bar and then purged using the high pressure product from column (2+). It is appreciated that the effluent from column (1+) in this step is fed to purge column (2-), which create an intermittent CH₄ and N₂ product at an intermediate pressure (24 bar). Column (1+) is then depressurized to a low pressure of 1.3 bar. This step takes approximately 15 seconds. Following this, column (1+) is purged for 45 seconds using the high pressure product of column (2+), or, alternatively, with stored intermittent product. Thus column (1+) can be purged without the use of a compressor. Column (1+) is then brought to a pressure of 13.8 bar for 5 seconds. This is accomplished by feeding column (1+) with the stream from columns (1-) and (2-) at high pressure. After this, the pressure is raised to 20 bar by equalizing the pressure of column (1+) with column (2-) for 15 seconds. Thus, the pressure of column (1+) is raised from 1.3 bar to 20 bar without the use of compressors. Finally, column (1+) is re-pressurized to 40 bar.

[0084] The cycle of column (2+) will now be described. The cycle of column (2+) commences as column (2+) is depressurized from 40 bar to 34.2 bar over a 30 second period. Simultaneously, column (2+) is connected to column (1-), such that the pressure of column (1-) can be increased while the pressure of column (2-) decreases. During this time period the pressure of column (2-) is brought from a pressure of 20 bar to 34.2 bar. After this step, the pressure of column (2+) is held for 30 seconds at 34.2 bar. This can be accomplished by closing

all of the valves of column (2+). Following this step, column (2+) is purged using the high pressure product from column (1+). It is appreciated that the effluent from column (2+) in this step is fed to purge column (1-), which creates an intermittent CH_4 and N_2 product at an intermediate pressure. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks. During this step the pressure is decreased to 13.8 bar.

[0085] The pressure of column (2+) is then further decreased to 1.3 bar. This step takes approximately 15 seconds. Next, column (2+) is purged at this low pressure for 45 seconds using the high pressure product of column (1+). After, column (2+) can be brought to a pressure of 13.8 bar for 5 seconds. This is accomplished by feeding column (2+) with the stream from columns (1-) and (2-) at high pressure and closing the other valve of column (2+). The pressure of column (2+) is further raised to 20 bar by equalizing the pressure of column (2+) with column (2-) for 15 seconds. Thus, column (2+) is purged and the pressure of column (2+) is raised from 1.3 bar to 20 bar without the use of compressors.

[0086] Column (2+) is next re-pressurized to 40 bar. Column (2+) then generates highly concentrated N_2 for 260 seconds during the high feed pressure step at 40 bar of pressure. Next, column (2+) is depressurized for 30 seconds and brought to a pressure of 34.2 bar. Simultaneously, column (2+) is connected to column (2-) such that the pressure of the two is equalized. During this time period the pressure of column (2-) is brought from a pressure of 20 bar to 34.2 bar. After this equalization step, the pressure of column (2+) is held for 15 seconds at 34.2 bar. This can be accomplished by closing all of the valves on column (2+).

[0087] Following this step, column (2+) is purged using the high pressure product from column (1+). It is appreciated that the effluent from column (2+) in this step is fed to purge column (2-), which creates an intermittent CH_4 and N_2 product at an intermediate pressure. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks. During this step the pressure is decreased to 13.8 bar.

[0088] Column (2+) is then depressurized to a low pressure of 1.3 bar, which takes approximately 15 seconds. Next, column (2+) is purged for 45 seconds using the high pressure feed of column (1+), or, alternatively, the stored intermittent product. Column (2+) is then brought to a pressure of 13.8 bar for 5 seconds. This is accomplished by feeding column (2+) with the stream from columns (1-) and (2-) at high pressure. After this, the pressure is raised to 20 bar by equalizing the pressure of column (2+) with column (1-) for 15 seconds. Thus, purging and increasing the pressure of column is accomplished without the use of compressors.

[0089] Column (2+) is next re-pressurized to 40 bar. After this, column (2+) is held at 40 bar and fed with a gaseous mixture of CH_4 and N_2 for 260 seconds. CH_4 is adsorbed in this column, while air with a high concentration of N_2 can be generated from the column.

[0090] The cycle of column (1-) will now be described. Column (1-) commences the cycle connected to column (2+) such that the pressure of column (1-) equalizes with the pressure of column (2-). In this step, the pressure of column (1-) is increased from 20 bar to 34.2 bar after 30 seconds. Next, the pressure of column (1-) is held at a constant 34.2 bar pressure for 30 seconds. Following this, column (1-) is fed with the purge effluent of (2+) (at 13.8 bar); this step creates intermittent CH_4 and N_2 products at 13.8 bar. During this step, the pressure of column (1-) is decreased from 34.2 bar to 20 bar over the course of 60 seconds. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks.

[0091] Column (1-) is next re-pressurized from 20 bar to 40 bar over a period of 65 seconds. After this, column (1-) is held at a pressure of 40 bar and fed with the high-pressure feed of the gaseous mixture of CH_4 and N_2 for 200 seconds. N_2 is adsorbed in this column and air with a high concentration of CH_4 is generated from the column. After this, column (1-) is depressurized to 36 bar over the course of 15 second. Column (1-) is then further depressurized to 24 bar and purged with the effluent of (2+) (at 24 bar); this step create intermittent CH_4 and N_2 products at 24 bar over the course of 60 seconds. This intermittent product may be stored and used as purge gas for low-pressure purges elsewhere in the cycle using appropriate storage tanks.

[0092] Column (1-) is again re-pressurized from 20 bar to 40 bar over a period of 65 seconds. After this, column (1-) is held at a pressure of 40 bar and fed with the high-pressure feed of the gaseous mixture of CH_4 and N_2 for 260 seconds. N_2 is adsorbed in this column and air with a high concentration of CH_4 is generated from the column. Column (1-) is then connected to column (2+), such that the pressure between the columns is equalized to an intermediate pressure, 20 bar. This takes 15 seconds. Next, the pressure of column (1-) is held at a constant 20 bar pressure for 60 seconds. This occurs until column (1-) is reconnected with column (1+) such that the pressure between the columns is equalized to an intermediate pressure, 34.2 bar for 30 seconds. Thus, the pressure of column (1-) is raised from 20 to 34.2 bar without the use of compressors.

[0093] Column (1-) is then re-pressurized from 34.2 bar to 40 bar over a period of 15 seconds. After this, column (1-) is held at a pressure of 40 bar and fed with the high-pressure feed of the gaseous mixture of CH_4 and N_2 for 140 seconds, as highly concentrated CH_4 is produced. Next Column (1-) is connected with column (1+) such that the pressure between the

two can equalize to 20 bar. This takes 15 seconds. Finally, the pressure in column (1-) is held for 60 seconds.

[0094] The cycle of column (2-) will now be described. Column (2-) commences the cycle in a high pressure feed step where column (2-) is held at a pressure of 40 bar and fed with the high-pressure feed of the gaseous mixture of CH_4 and N_2 for 185 seconds. N_2 is adsorbed in this column and air with a high concentration of CH_4 is generated from the column. Column (2-) is then connected to column (2+), such that the pressure between the columns is equalized to an intermediate pressure, 20 bar. This takes 15 seconds. Next, the pressure of column (2-) is held at a constant 20 bar pressure for 60 seconds. This occurs until the column (2-) is reconnected with column (1+) such that the pressure between the columns is equalized to an intermediate pressure, 34.2 bar for 30 seconds. Thus, the pressure of column (2-) is raised from 20 to 34.2 bar without the use of compressors.

[0095] Column (2-) is then re-pressurized from 34.2 bar to 40 bar over a period of 15 seconds. After this, column (2-) is held at a pressure of 40 bar and fed with the high-pressure feed of the gaseous mixture of CH_4 and N_2 for 140 seconds. N_2 is adsorbed in this column and air with a high concentration of CH_4 is generated from the column. Column (2-) is then connected to column (1+), such that the pressure between the columns is equalized to an intermediate pressure, 20 bar. This takes 15 seconds. Next, the pressure of column (2-) is held at a constant 34.2 bar pressure for 60 seconds. This occurs until column (2-) is reconnected with column (2+), such that the pressure between the columns is equalized to an intermediate pressure of 34.2 bar for 30 seconds. This is briefly followed by a hold period, where the pressure remains unchanged in column (2-) for 34.2 seconds. Following this, column (2-) is purged with the effluent of (2+) (at 13.8 bar). This step creates intermittent CH_4 and N_2 products. During this step, the pressure of column (2-) is decreased from 34.2 bar to 13.8 bar over the course of 60 seconds.

[0096] Column (2-) is next re-pressurized from 13.8 bar to 40 bar over a period of 65 seconds. After this, column (2-) is held at a pressure of 40 bar and fed with the high-pressure feed of the gaseous mixture of CH_4 and N_2 for 200 seconds. N_2 is adsorbed in this column and air with a high concentration of CH_4 is generated from the column. Following this step, column (2-) is connected to columns (1-) and the pressure is equalized between them to 34.2 bar. This takes 15 seconds. Following this, column (2-) is depressurized to 24 bar and purged using the effluent of column (1-) at 24 bar. This takes 60 seconds and creates an intermittent CH_4 and N_2 product.

[0097] Column (2-) is then re-pressurized from 20 bar to 40 bar over a period of 65 seconds. After this, column (1-) is held at a pressure of 40 bar and fed with the high-pressure feed of the gaseous mixture of CH₄ and N₂ for 40 seconds. N₂ is adsorbed in this column and air with a high concentration of CH₄ is generated from the column. As can be seen in FIG. 4 and just described, columns (2-) and (1-) are connected in such a way that a high concentration of CH₄ is continuously produced throughout the cycle.

[0098] The embodiments and examples described herein are merely illustrative, as numerous other embodiments may be implemented without departing from the spirit and scope of the exemplary embodiments of the present application. Moreover, while certain features of the application may be shown on only certain embodiments or configurations, these features may be exchanged, added, and removed from and between the various embodiments or configurations while remaining within the scope of the application. Likewise, methods described and disclosed may also be performed in various sequences, with some or all of the disclosed steps being performed in a different order than described while still remaining within the spirit and scope of the present application.

CLAIMS:

1. A pressure swing adsorption system for the fractionation of a multicomponent fluid comprising a first adsorbent column selective for a first component, a second adsorbent column selective for the first component, a third adsorbent column selective for a second component, a fourth adsorbent column selective for the second component;

wherein said columns are fed with the multicomponent fluid while pressurized at a high pressure during at least one phase of a pressure swing adsorption cycle that enables a continuous supply of both a highly concentrated first component and a highly a concentrated second component to be recovered throughout the pressure swing adsorption cycle;

wherein said columns are depressurized from the high pressure to a first intermediate pressure during at least one phase of the pressure swing adsorption cycle;

wherein said columns are purged while pressurized at a second intermediate pressure during at least one phase of the pressure swing adsorption cycle and an intermittent supply of first and second component is created;

wherein said columns are depressurized from a third intermediate pressure to a low pressure during at least one phase of the pressure swing adsorption cycle;

wherein said columns are purged while pressurized at the low pressure during at least one phase of the pressure swing adsorption cycle by using an effluent from a column selective for a different component or from said intermittent supply;

wherein said columns are re-pressurized to the high pressure during at least one phase of the pressure swing adsorption cycle.

2. The pressure swing adsorption system of claim 1, wherein the first, second and third intermediate pressures are the same.

3. The pressure swing adsorption system of claim 1, wherein the system operates without the use of a compressor.

4. The pressure swing adsorption system of claim 1, wherein an adsorbent column selective for a first type of component is connected to a column selected for a second type of component during at least one phase of the pressure swing adsorption cycle to enable pressure equalizations.

5. The pressure swing adsorption system of claim 4, wherein the four adsorption columns are placed in a partitioned vessel with common walls.

6. The pressure swing adsorption system of claim 4, wherein at least two of the adsorption columns are placed in a partitioned vessel with common walls.

7. The pressure swing adsorption system of claim 4, wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for carbon dioxide.
8. The pressure swing adsorption system of claim 4, wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for methane.
9. The pressure adsorption system of claim 7, wherein
the adsorbent selective for nitrogen is selected from the group consisting of 4A or NaA zeolite;
the adsorbent selective for carbon dioxide is selected from the group consisting of activated carbons, aluminas, metal organic framework, mixed metal oxides, hydrotalcites or combinations thereof.
10. The pressure swing adsorption system of claim 8, wherein
the adsorbent selective for methane is selected from the group consisting of H/Na mordenite, activated carbons, aluminas, metal organic framework, mixed metal oxides, hydrotalcites or combinations thereof;
the adsorbent selective for nitrogen is selected from the group consisting of clinoptilolite, rho-zeolite, ITQ or carbon molecular sieve or combinations thereof.
11. The pressure swing adsorption system of claim 8, wherein
the ratio of the time that the adsorbent column selective for methane is fed with the multicomponent fluid while pressurized at a high pressure to total cycle time is between 0.4 and 0.6;
the ratio of the time that the adsorbent column selective for methane is purged to the total cycle time is between 0.15 and 0.25;
the ratio of the time that the adsorbent column selective for nitrogen is fed with the multicomponent fluid while pressurized at a high pressure to total cycle time is between 0.4 and 0.6;
the ratio of the time that the adsorbent column selective for nitrogen is purged to the total cycle time is between 0.08 and 0.16;
12. The pressure swing adsorption system of claim 4, wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for oxygen.
13. The pressure swing adsorption system of claim 4, wherein two columns contain an adsorbent selective for propane and two columns contain an adsorbent selective for propylene.
14. The pressure swing adsorption system of claim 4, wherein two columns contain an adsorbent selective for ethane and two columns contain an adsorbent selective for ethylene.

15. A method for fractionation of a multicomponent stream having first and second components and for providing a continuous purified supply of said multicomponent stream using a pressure swing adsorption system having first adsorption columns selective for a first component and second adsorption columns selective for a second component, which comprises of steps of:
- a) feeding an adsorption column, selective for the first component, at a high pressure with a stream of the multicomponent gas and generating a product comprising the second component;
 - b) depressurizing said column to a first intermediate pressure, which is between the high pressure and a low pressure;
 - c) purging said column at a second intermediate pressure by using one of the product of the second component or from the purge effluent of a column selective for the second component;
 - d) depressurizing said column from a third intermediate pressure to the low pressure;
 - e) purging said column at the low pressure by using product of second type of component; and
 - f) increasing the pressure of said column from to the high pressure
16. The method of claim 14, wherein said column is connected to another column selective for the second component to enable to pressure between said column and another column to equalize without the use of a compressor.
17. The method of claim 16 wherein the first, second and third intermediate pressures are the same.
18. The method of claim 16 wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for carbon dioxide.
19. The method of claim 16, wherein the four adsorption columns are placed in a partitioned vessel with common walls.
20. The method of claim 16, wherein at least two of the adsorption columns are placed in a partitioned vessel with common walls.
21. The method of claim 16, wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for carbon dioxide.
22. The method of claim 16, wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for methane.
23. The method of claim 21, wherein

the adsorbent selective for nitrogen is selected from the group consisting of 4A or NaA zeolite;

the adsorbent selective for carbon dioxide is selected from the group consisting of activated carbons, aluminas, metal organic framework, mixed metal oxides, hydrotalcites or combinations thereof.

24. The method of claim 22, wherein

the adsorbent selective for methane is selected from the group consisting of H/Na mordenite, activated carbons, aluminas, metal organic framework, mixed metal oxides, hydrotalcites or combinations thereof;

the adsorbent selective for nitrogen is selected from the group consisting of clinoptilolite, rho-zeolite, ITQ or carbon molecular sieve or combinations thereof.

25. The method of claim 22, wherein

the ratio of the time that the adsorbent column selective for methane is fed with the multicomponent fluid while pressurized at a high pressure to total cycle time is between 0.4 and 0.6;

the ratio of the time that the adsorbent column selective for methane is purged to the total cycle time is between 0.15 and 0.25;

the ratio of the time that the adsorbent column selective for nitrogen is fed with the multicomponent fluid while pressurized at a high pressure to total cycle time is between 0.4 and 0.6;

the ratio of the time that the adsorbent column selective for nitrogen is purged to the total cycle time is between 0.08 and 0.16;

26. The method of claim 16, wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for oxygen.

27. The method of claim 16, wherein two columns contain an adsorbent selective for propane and two columns contain an adsorbent selective for propylene.

28. The method of claim 16, wherein two columns contain an adsorbent selective for ethane and two columns contain an adsorbent selective for ethylene.

29. A method for fractionation of a multicomponent stream having first and second components and for providing a continuous purified supply of said multicomponent stream using a thermal swing adsorption system, wherein first adsorption columns are used that are selective for the first component and second adsorption columns are used that are selective for the second component, which comprises of steps of:

- a) feeding an adsorption column, selective for the first component, at a first high temperature with a stream of the multicomponent stream and generating a product comprising the second component;
 - b) decreasing the temperature of said column by coming to thermal equilibrium with another column which is at a first low temperature;
 - c) purging said column at a first intermediate temperature by using one of the product comprising the second component or the purge effluent of a column selective for the second component;
 - d) purging said column at a second low temperature by using the product comprising the second component; and
 - e) increasing the temperature of said column from a third low temperature to a third high temperature, by coming to thermal equilibrium with another column which is at higher temperature than said column
30. The method of claim 29, wherein the system operates without the use of a compressor.
- 31 The method of claim 29, wherein the first, second, and third low temperatures are the same.
32. The method of claim 29, wherein the first, second, and third high temperatures are the same.
33. The method of claim 29, wherein the four adsorption columns are placed in a partitioned vessel with common walls; wherein
the common walls are thermally conductive.
34. The method of claim 29, wherein at least two of the adsorption columns are placed in a partitioned vessel with common walls; wherein
the common walls are thermally conductive.
35. The method of claim 29 wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for carbon dioxide.
36. The method of claim 29, wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for methane.
37. The method of claim 29, wherein two columns contain an adsorbent selective for nitrogen and two columns contain an adsorbent selective for oxygen.
38. The method of claim 29, wherein two columns contain an adsorbent selective for propane and two columns contain an adsorbent selective for propylene.
39. The method of claim 29, wherein two columns contain an adsorbent selective for ethane and two columns contain an adsorbent selective for ethylene.

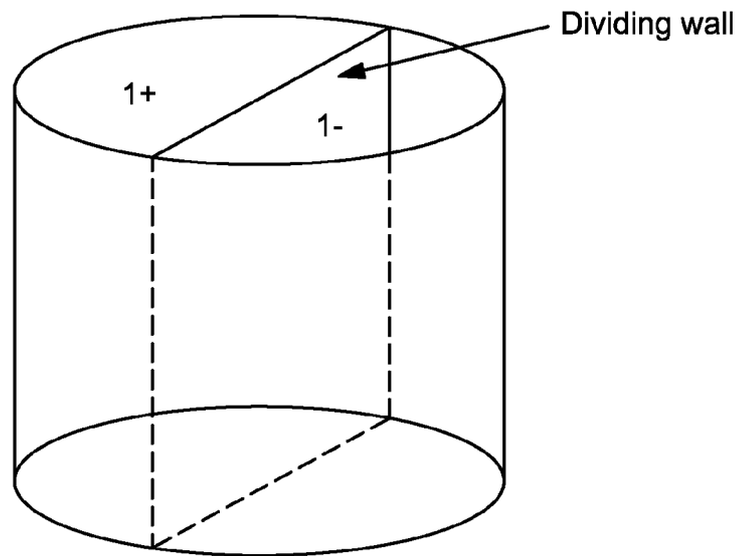


FIG. 2

3/5

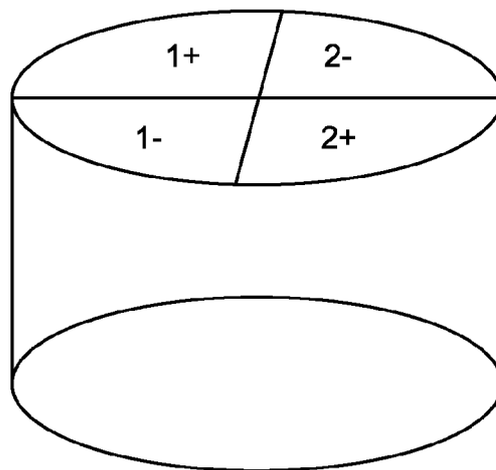


FIG. 3

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US2016/059916

Box No. II Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. Claims Nos.:
because they relate to subject matter not required to be searched by this Authority, namely:

2. Claims Nos.:
because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

3. Claims Nos.:
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box No. III Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.

2. As all searchable claims could be searched without effort justifying an additional fees, this Authority did not invite payment of additional fees.

3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos. :

4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos. :

1-28

Remark on Protest

- The additional search fees were accompanied by the applicant's protest and, where applicable, the payment of a protest fee.
- The additional search fees were accompanied by the applicant's protest but the applicable protest fee was not paid within the time limit specified in the invitation.
- No protest accompanied the payment of additional search fees.

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2016/059916

A. CLASSIFICATION OF SUBJECT MATTER
INV. B01D53/047
ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
B01D

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-Internal , WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 4 744 803 A (KNAEBEL KENT S [US]) 17 May 1988 (1988-05-17) cited in the applicati on figures 1-3	1-28
A	----- EP 1 022 045 A2 (BOC GROUP INC [US]) 26 July 2000 (2000-07-26) paragraphs [0046] , [0048] ; figure 9	1, 15
A	----- Wo 2008/151913 AI (AI R LIQUIDE [FR] ; SCHNEIDER MARTINE [FR] ; DE SOUZA GUI LLAUME [FR] ; ALB) 18 December 2008 (2008-12-18) figure 2	1, 15
A	----- EP 0 212 494 AI (LINDE AG [DE]) 4 March 1987 (1987-03-04) figure 1	1, 15
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Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents :

"A" document defining the general state of the art which is not considered to be of particular relevance

"E" earlier application or patent but published on or after the international filing date

"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

"O" document referring to an oral disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search

30 January 2017

Date of mailing of the international search report

30/03/2017

Name and mailing address of the ISA/
European Patent Office, P.B. 5818 Patentlaan 2
NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040,
Fax: (+31-70) 340-3016

Authorized officer

Bergt, Thomas

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2016/059916

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 4 299 596 A (BENKMANN CHRISTIAN) 10 November 1981 (1981-11-10) figures 4-5 -----	1, 15

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No PCT/US2016/059916
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FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. claims: 1-28

The common technical features present in all independent claims are: Adsorption from a mixture in complementary (different) adsorbents selective for two different components of the mixture.
PSA method/system with intermediate pressure purge.

2. claims: 29-39

The common technical features present in all independent claims are: Adsorption from a mixture in complementary (different) adsorbents selective for two different components of the mixture.
TSA method with intermediate/low temperature purge step(s) .
