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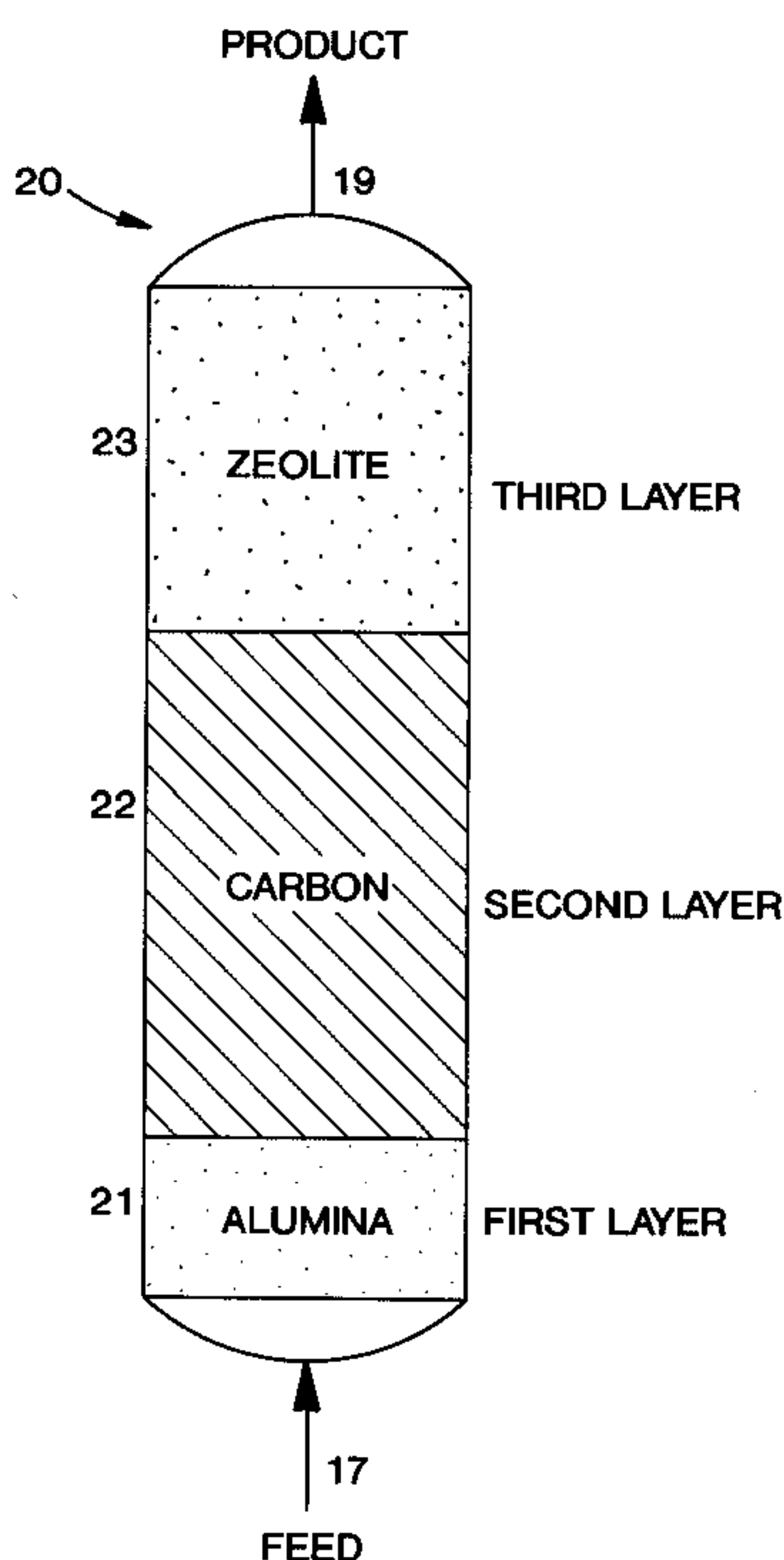
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(54) **PROCEDE D'ADSORPTION MODULEE EN PRESSION POUR
LA PRODUCTION D'HYDROGENE**

(54) **PRESSURE SWING ADSORPTION PROCESS FOR THE
PRODUCTION OF HYDROGEN**



(57) This invention provides a pressure swing adsorption process for purifying a synthesis gas stream containing from 60 to 90 mole % hydrogen and impurities such as CO₂, CH₄, N₂, and CO. The PSA process of the invention further provides a method of adsorbing substantially all of the nitrogen and other contaminants from the feed gas stream; wherein the feed stream is passed at superatmospheric pressure through a plurality of adsorbent beds and each adsorbent bed contains at least a CaX, LiA, LiX or calcium containing mixed cation zeolite having a SiO₂/Al₂O₃ mole ratio of 2.0-2.5. Such process involves sequentially pressurizing, depressurizing, purging and repressurizing the adsorbent beds with product hydrogen, and recovering product hydrogen in purities of 99.9% or greater from the beds.



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ABSTRACT OF THE DISCLOSURE

This invention provides a pressure swing adsorption process for purifying a synthesis gas stream containing from 60 to 90 mole % hydrogen and impurities such as CO₂, CH₄, N₂, and CO. The PSA process of the invention further provides a method of adsorbing substantially all of the nitrogen and other contaminants from the feed gas stream; wherein the feed stream is passed at superatmospheric pressure through a plurality of adsorbent beds and each adsorbent bed contains at least a CaX, LiA, LiX or calcium containing mixed cation zeolite having a SiO₂/Al₂O₃ mole ratio of 2.0-2.5. Such process involves sequentially pressurizing, depressurizing, purging and repressurizing the adsorbent beds with product hydrogen, and recovering product hydrogen in purities of 99.9% or greater from the beds.

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PRESSURE SWING ADSORPTION
PROCESS FOR THE PRODUCTION OF HYDROGEN

FIELD OF THE INVENTION

This invention relates to a pressure swing adsorption (PSA) process for purifying impure gas streams containing more than 50 mole % hydrogen, and more particularly to such a process for the production of high purity hydrogen from various hydrogen-containing feed mixtures such as synthesis gas. The improved process provides higher hydrogen recovery and lower adsorbent inventory as compared with previously known PSA processes for hydrogen production.

BACKGROUND OF THE INVENTION

The need for high purity (>99.9%) hydrogen is growing in the chemical process industries, e.g. in steel annealing, silicon manufacturing, hydrogenation of fats and oils, glass making, hydrocracking, methanol production, the production of oxo alcohols and isomerization processes. This growing demand requires the development of highly efficient separation processes for H₂ production from various feed mixtures. In order to obtain highly efficient PSA separation processes, both the capital and operating costs of the PSA system must be reduced.

One way of reducing PSA system cost is to decrease the adsorbent inventory and number of beds in the PSA process. In addition, further improvements may be possible using advanced cycles and adsorbents in the PSA process. However, 1-12 feed gas contains several contaminants, e.g. CO₂ (20% to 25%) and minor amounts

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of H₂O (<0.5%), CH₄ (<3%), CO (<1%) and N₂ (<1%). Such a variety of adsorbates at widely varying compositions, combined with the high purity (>99.9%) requirement for H₂, presents a significant challenge to efficient selection, configuration and amount of adsorbents in each layer of the bed to achieve an efficient H₂-PSA process.

There are a variety of known processes for producing hydrogen. For example, Figure 1 of the accompanying drawings shows the steam reforming of natural gas or naphtha wherein a feedstock, e.g. a natural gas stream 11, is compressed and fed to a purification unit 12 to remove sulfur compounds. The desulfurized feed is then mixed with superheated steam and fed to a reformer 13 to produce primarily H₂ and CO. The effluent stream from the reformer is sent to a heat recovery unit 14, then to a shift converter 15 to obtain additional H₂. The effluent from the shift converter is cooled and recovered in unit 16. The effluent, synthesis gas stream 17, having on a dry basis a composition of about 74.03% H₂, 22.54% CO₂, 0.36% CO, 2.16% CH₄, and 0.91% N₂ is then routed to a PSA purification system 18 to produce a high purity hydrogen product stream 19.

Representative prior art PSA processes for hydrogen purification include the following: (1) Wagner, U.S. Pat. No. 3,430,418, (2) Batta, U.S. Pat. No. 3,564,816, (3) Sircar et al., U.S. Pat. No. 4,077,779, (4) Fuderer et al., U.S. Pat. No. 4,553,981, (5) Fong et al, U.S. Pat. No. 5,152,975 and (6) Kapoor et al., U.S. Pat. No. 5,538,706.

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The adsorbers in hydrogen PSA processes have been conceptually divided into multiple zones, depending upon the particular contaminants to be removed in the successive zones. For example, in Wagner (U.S. Pat. No. 3,430,418) a combination of two types of adsorbents is used, i.e. activated carbon for the removal of H₂O and CO₂, and calcium zeolite A for removal of CO and CH₄ (see Example 1). The Wagner patent describes an eight-step PSA cycle for hydrogen purification. At least four beds are used in the process; following the bed-to-bed equalization step, each bed undergoes a co-current depressurization step prior to countercurrent blowdown to recover void space gas for purging of another bed.

Batta (U.S. Pat. No. 3,564,816) describes a twelve-step PSA cycle using at least four adsorbent beds and two pressure equalization stages for separating hydrogen-containing gas mixtures contaminated with H₂O, CO₂, CH₄ and CO produced in the steam reforming of natural gas. In the Batta process, a co-current depressurization step follows the first bed-to-bed equalization step to recover void space gas for purging of another bed. A second bed-to-bed equalization step is used prior to the countercurrent blowdown step in the PSA cycle.

Sircar, (U.S. Pat. No. 4,171,206), discloses a PSA process in which a crude hydrogen stream (such as the gaseous effluent from a shift converter of a hydrocarbon reforming plant) flows through a first bed of activated carbon (effective for the removal of CO₂), and then through a second bed of 5A zeolite (effective

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for the removal of dilute impurities such as CH₄ and/or CO) to produce high purity (>99.9%) hydrogen.

Golden et al, (U.S. Pat. No. 4,957,514), discloses the purification of hydrogen using a barium-exchanged Type X zeolite to remove CO, CH₄ and N₂ contaminants. According to Golden, the preferred BaX zeolite is one in which 60 to 100% of the sodium cations of a NaX zeolite have been replaced by barium cations. Golden compares the adsorbent requirements using BaX (96% Ba, 4% Na), CaX (98% Ca, 2% Na), Ca/SrX (50% Ca, 50% Sr), and commercial 5A zeolites in hydrogen purification processes. For a given feed flow rate and H₂ purity, the quantity of zeolite required in the hydrogen purification process is lowest when BaX is used. Also, Golden ranks the adsorbents in the order BaX > Ba/SrX > 5A > SrX > Ca/SrX > CaX for CO or CH₄ adsorption. In particular, CaX is ranked the lowest with respect to the removal of CO and CH₄ impurities.

Scharpf et al, (U.S. Pat. No. 5,294,247), discloses a vacuum PSA process for recovering hydrogen from dilute refinery off gases, preferably containing less than 60% hydrogen. The patent discloses the use of six adsorbent beds, each of which contains a layer of activated carbon, a layer of 13X zeolite, a layer of 5A zeolite and a layer of CaA zeolite or calcium exchanged X zeolite. This four-layer arrangement is described as useful for the removal of large feed concentrations (>1%) of CO and CO₂.

More recently, Bomard et al in International Patent Application W097/45363, disclosed a method for separating hydrogen from a gas mixture that contains CO

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and other impurities such as CO₂ and hydrocarbons. In the Bomard application, the feed mixture is passed into a first selective adsorbent (e.g., activated carbon) to remove CO₂ and hydrocarbons. It is then contacted by a second adsorbent, a faujasite-type zeolite with at least 80% lithium exchange, to remove primarily CO impurity and to produce high purity hydrogen. In addition, a third adsorbent (5A zeolite) may be placed between the first and second adsorbents to remove nitrogen if N₂ is also present in the feed mixture.

It is also known to utilize PSA processes for the selective adsorption of N₂ from air to obtain O₂-enriched gases. Berlin, (U.S. Pat. No. 3,313,091), describes the use of strontium-substituted type X zeolites in such a process and stipulates that Ca²⁺, Sr²⁺, and Ag⁺ are the preferred exchanged cations with Sr²⁺ being most desirable. In the case of Type A zeolite, Ca²⁺, Mg²⁺ and Ag⁺ are preferred.

Coe et al, (U.S. Pat. No. 4,481,018), also discloses PSA air separation processes involving N₂ and O₂ separations using calcium-exchanged forms of zeolite X, and showed enhanced PSA process performance with increasing calcium content. However, Chao (U.S. Pat. Nos. 5,698,013 and 5,454,857) discloses for calcium-exchanged forms of zeolite X a peak performance in air separation below maximum calcium content. In particular, the peak performance occurs when the degree of calcium exchange is in the range of 60 to 89 equivalent percent, and the SiO₂/AlO₂O₃ molar ratio is in the range of 2.0 to 2.4.

The operating conditions used in H₂ PSA processes are distinct from those used in PSA processes for O₂

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production from air. The adsorption pressure is typically less than 2.5 bars in VPSA air separation, while the adsorption pressure is 5 to 20 bars in H₂ PSA.

The amount of N₂ in the feed streams of these two processes is significantly different, e.g. the N₂ fraction in air is about 78 mole %; whereas, in hydrogen purification, the proportion of N₂ in the feed stream is typically less than 1-3 mole %. Consequently, the N₂-selective adsorbents disclosed in the aforementioned references for air separation must perform in different N₂ partial pressure regions of the isotherm than those required for the function of purification in H₂ PSA processes. Furthermore, the N₂ differential loadings in O₂ VPSA and H₂ PSA processes are quite different for a given N₂-selective adsorbent (e.g., zeolite). Additional complications in the design of H₂PSA processes result from the competitive adsorption and diffusion rates of each adsorbate in the mixture. In selecting improved adsorbents and designing H₂ PSA processes, due consideration must be given to all of the foregoing issues.

The adsorbent typically used heretofore in H₂ PSA processes is 5A zeolite, obtained by base exchange of Ca (about 75%) with the sodium ion present in 4A zeolite.

It is among the objects of the present invention to provide an improved PSA process for the production of hydrogen from an impure gas stream containing more than 50 mole % hydrogen, such improvements realized as increased hydrogen recovery, reduced adsorbent and lower capital and operating costs. Other objects and

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advantages of the invention will be apparent from the following description taken in connection with the accompanying drawing.

SUMMARY OF THE INVENTION

In accordance with the present invention, an improved PSA process is provided for purifying a gas stream containing more than about 50 mole % hydrogen, which comprises adsorbing substantially all of the nitrogen in such stream by passing it at superatmospheric pressure through an adsorbent bed containing CaX, LiA or LiX-type zeolite adsorbent having SiO₂/Al₂O₃ mole ratio within the range of 2.0-2.5, and recovering purified hydrogen as product from the adsorbent bed. By providing additional adsorbent layers to remove other impurities such as H₂O, CO₂, CH₄ and CO upstream of the CaX, LiA or LiX zeolite adsorbent, a high purity (≥99.9%) hydrogen product is recovered.

In accordance with a further feature of the invention, substantially all of the CO₂ is first removed, i.e. leaving less than about 0.15 mole %, preferably less than about 0.10 mole % CO₂ for subsequent passage into the zeolite layer where this residual CO₂ is removed along with the N₂. Configuring the adsorbent bed according to this contaminant removal strategy increases H₂ recovery and reduces the quantity of zeolite required for N₂ removal compared to prior art PSA processes for H₂ production.

The preferred adsorbents utilized in the process of the present invention are CaX zeolites, most desirably CaX (2.0), a zeolite of the faujasite type

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exchanged at least 90% with calcium and having a $\text{SiO}_2/\text{Al}_2\text{O}_3$ molar ratio of 2.0. CaX (2.0) allows the processing of more feed gas per unit weight of adsorbent at a given P/F (purge to feed) ratio than other N_2 -selective adsorbents. Accordingly, by using CaX (2.0) the amount of the adsorbent required for removing N_2 may be significantly reduced, i.e. reducing the bed size factor (BSF). Moreover, this reduction in bed size factor results in lower H_2 losses during regeneration of the adsorbent bed. This in turn leads to higher hydrogen recovery than may be obtained with other N_2 -selective adsorbents.

BRIEF DESCRIPTION OF THE DRAWINGS

In the accompanying drawing:

Figure 1 is a schematic of the prior art technique for the production of hydrogen by the steam reforming of natural gas;

Figure 2 is a comparison of the N_2 adsorption isotherms of CaX (2.0) and other N_2 -selective zeolite adsorbents, at 300°K ;

Figure 3 is a comparison of the differential N_2 loading of CaX (2.0) with other adsorbents;

Figure 4 is a comparison of the CO_2 adsorption isotherms of CaX (2.0) and other N_2 -selective adsorbents, at 300°K ;

Figure 5 is a schematic drawing of a PSA adsorption bed of the present invention;

Figure 6 is a schematic drawing of a four-bed system for carrying out the

PSA process of the invention;

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Figure 7 is a graphic depiction of the representative bed pressure profile during one complete PSA cycle in the preferred embodiment of the invention described herein;

Figure 8 is a graphic illustration of the representative gaseous component mole fractions in an adsorbent bed in the embodiment described, at the conclusion of the high pressure adsorption step therein;

Figure 9 is a graphic comparison of the computer simulated PSA process performance for the system of Figure 5, utilizing each of the CaX (2.0), 5A, LiX and VSA6 zeolite adsorbents in the N₂-selective layer;

Figure 10 is a graphic comparison of the quantity of CaX (2.0), 5A, LiX and VSA6 zeolite adsorbents required in the illustrated four bed PSA process; and

Figure 11 is a graphic comparison of differential loading of N₂ on CaX (2.0), 5A, LiX and VSA6 zeolites.

DETAILED DESCRIPTION OF THE INVENTION

As indicated hereinabove, the PSA process of the present invention provides improved recovery of high purity (greater than 99.9%) hydrogen from gas streams containing more than about 50 mole % H₂, and preferably from about 60 to 90 mole hydrogen. The process is particularly applicable to the purification of synthesis gas produced in feed streams recovered in reforming processes such as outlined in Figure 1 of the drawing. Such streams may contain from 60 to 90 mole % hydrogen with impurities that include CO₂, H₂O, CH₄, N₂ and CO.

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Desirably, the purification is affected by passing the stream at superatmospheric pressure through a plurality of adsorbent beds each of which contains at least one adsorbent layer containing CaX, LiA or LiX zeolite having a $\text{SiO}_2/\text{Al}_2\text{O}_3$ ratio within the range of 2.0-2.5 for adsorbing substantially all of the N_2 from the stream. The process includes the steps of sequentially pressurizing, depressurizing, purging and repressurizing the adsorbent beds with product hydrogen to obtain the desired hydrogen product in purities of 99.9% or greater as unadsorbed effluent from the beds.

As indicated herein above, the preferred CaX zeolite adsorbent utilized in the practice of this invention is CaX (2.0), an adsorbent which has been found to provide superior nitrogen adsorption as compared with other N_2 -selective adsorbents. Other useful Ca-exchanged zeolites may be prepared from naturally occurring crystalline zeolite molecular sieves such as chabazite, erionite, clinoptilolite and faujasite. Alternatively, the CaX zeolites useful herein include mixed cation (e.g. Ca^{2+} and Na^+) zeolites such as VSA-6 developed by UOP of Des Plaines, IL with 74% Ca^{2+} and a $\text{SiO}_2/\text{Al}_2\text{O}_3$ ratio of 2.3. In general, by "mixed cations" we mean an adsorbent containing at least two different cations. Such adsorbents would further include for example LiSrX, CaLiX, CaNaX, etc.

LiA and LiX zeolites having $\text{SiO}_2/\text{Al}_2\text{O}_3$ ratios within the range of 2.0-2.5 are useful in the processes described above. Other adsorbents displaying improved performance include mixed lithium/alkaline earth metal Type A and Type X zeolites having $\text{SiO}_2/\text{Al}_2\text{O}_3$ molar ratios in the range of 2.0-2.5 such as CaLiX (2.3),

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having calcium contents of 15-30% (see Chao et al, (U.S. Pat. Nos. 5,413,625; 5,174,979; 5,698,013; 5,454,857 and 4,859,217). The zeolite disclosures of the foregoing patents are incorporated by reference herein.

The superiority of CaX (2.0) as an adsorbent for nitrogen is illustrated in Figures 2 and 3. Figure 2 compares the nitrogen adsorption isotherms at 300°K of CaX (2.0) with other zeolites; whereas, Figure 3 compares the N₂ differential loadings on several adsorbents. The N₂ differential loading for each adsorbent in Figure 3 was determined as the difference in loading of N₂ on the adsorbent between the end of adsorption and desorption steps in the cycle. The pressure, temperature and N₂ composition at the end of adsorption and desorption steps used in the calculation are (11.7bar, 306°K, Y_{N₂} = 0.008) and (1.36bar, 306°K, Y_{N₂} = 0.025), respectively. Pure component isotherm data were used in conjunction with a multicomponent isotherm model to determine the loadings at the conditions indicated.

Figure 2 shows that CaX (2.0) exhibits substantially greater N₂ adsorption than 5A and VSA6 across a broad range of adsorption partial pressures. It is further evident from Figure 3 that CaX (2.0), LiX (2.3), VSA6, LiX (2.0), CaLiX (2.3), and the CaX(2.0)/13X mix exhibit superior differential N₂ loading as compared with 13X, 5A, NaY, activated alumina (A201) and activated carbon.

Figure 4 of the drawing illustrates the comparative amount of CO₂ adsorbed by various zeolites at, 300°K. CO₂ is strongly adsorbed on CaX (2.0) at

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CO₂ partial pressures in excess of about 2 atmospheres - CaX solely exceeded by LiX in this regard. When a substantial amount of CO₂ is adsorbed on these N₂-selective zeolites regeneration by pressure swing adsorption is extremely difficult, i.e. at the typical operating conditions of H₂ PSA processes. When CO₂ is coadsorbed with N₂, adsorption capacity for N₂ and other trace impurities is diminished, resulting in degradation of PSA process performance and hydrogen purity. Accordingly as indicated above, it is desirable to adsorb other impurities, e.g. CO₂, H₂O, CH₄ and CO, from the feed gas stream prior to their passing through the CaX, LiA or LiX zeolite layer. In particular, it is important to reduce the CO₂ content of the gas stream before such stream reaches the zeolite to less than about 0.15 mole %, preferably to less than about 0.05 to 0.10 mole %, thereof.

It is preferred to utilize one or more adsorbent beds incorporating multiple layers for removal of the foregoing impurities in the practice of the present invention. One preferred layered configuration is illustrated in Figure 5 showing adsorbent bed embodiment 20. Bed 20 includes a first alumina layer 21 for removing H₂O from the feed gas stream, i.e. the synthesis gas stream 17 of the reforming technique illustrated in Figure 1. Layer 21 is followed by an activated carbon layer 22 for removing CO₂ from the feed gas stream to concentrations less than about 0.15 mole %. Finally, a CaX, LiA or LiX adsorbent layer 23 is located at the product end of the bed for removing N₂, thereby producing the desired high purity H₂ product stream 19. One skilled in the art will

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recognize that other adsorbents may be substituted for alumina for H₂O adsorption in layer 21, e.g. zeolites, silica gel, and that other CO₂-selective adsorbents may be substituted for activated carbon in layer 22 of bed 20.

The H₂ PSA process of the invention is suitably carried out at conditions previously utilized in the art, e.g. as described in U.S. Pat. No. 3,564,816, the process parameters of which are incorporated by reference herein. Thus, N₂ adsorption in the CaX, LiA or LiX adsorbent layer may be carried out at temperatures from about 250 to 350°K, and at total pressures from about 5 to 20 bars. The adsorbent layer may thereafter be depressurized and purged at temperatures of from about 250 to 350°K and at total pressures from about 0.50 to 1.70 bars. Specific temperature, pressure and other operating conditions utilized in the H₂ PSA system incorporating the CaX, LiA or LiX adsorbent in accordance with the present invention will depend upon the design of the particular PSA system. The choice of operating conditions will also depend upon the adsorbents selected for the other layers in the bed, the feed gas composition and flow rate and other parameters associated with the integration of the PSA unit with the other unit operations shown in Figure 1.

Examples

The present invention will be more fully described in connection with the following computer-simulated embodiments employing the four-bed system illustrated in Figure 6 of the drawings, each bed of which utilizes

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alumina, activated carbon and an N₂-selective adsorbent arranged in layers as shown below. The bed pressure and axial gas phase concentration profiles are shown in Figures 7 and 8, respectively. The valve switching logic, the time interval, and step sequence of the PSA cycle are shown in Table 1 below.

The sequence for the twelve step illustrative cycle is as follows:

Step 1 (AD1): Bed 1 (B1) is in the first adsorption step (AD1) at 11.72 bars, while bed 2 (B2) is undergoing countercurrent blowdown (BD), bed 3 (B3) is undergoing the first equalization falling step (EQ1DN), and bed 4 (B4) is undergoing the second pressure equalization rising step (EQ2UP).

Step 2 (AD2): Bed 1 is in the second adsorption step (AD2) and is also supplying product gas to bed 4 that is undergoing the first product pressurization (PP1) step. During the same time, beds 2, 3 and 4 are undergoing purge, concurrent depressurization and first product pressurization, respectively.

Step 3 (AD3): Bed 1 is in the third adsorption step (AD3), and is also supplying product gas to bed 4 that is undergoing the second product pressurization (PP2) step. During the same time period, beds 2, 3 and 4 are undergoing the first equalization rising step (EQ1UP), second equalization falling step (EQ2DN), and second product pressurization step (PP2), respectively.

Step 4 (EQ1DN): Bed 1 is undergoing the first equalization falling step (EQ1DN), while bed 2 receives the gas from bed 1 and is undergoing the second equalization rising step (EQ2UP). Beds 3 and 4 are now

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undergoing blowdown (BD) and the first adsorption step (ADI), respectively.

Step 5 (PPG): Bed 1 is undergoing a cocurrent depressurization step to provide purge gas (PPG) to bed 3, while beds 2 and 4 are undergoing the first product pressurization (PP1) and the second adsorption step (AD2), respectively.

Step 6 (EQ2DN): Bed 1 undergoes a second equalization falling step (EQ2DN) by sending low pressure equalization gas to bed 3 that is undergoing the first equalization rising step (EQ1UP). Beds 2 and 4 are undergoing the second product pressurization (PP2) and third adsorption step, respectively.

Step 7 (BD): Beds 1 and 2 undergo the countercurrent blowdown (BD) and first adsorption (AD1) step, respectively. During this time beds 3 and 4 are undergoing bed-to-bed equalization, i.e., beds, 3 and 4 are undergoing the second equalization rising (EQ2UP) and first equalization falling (EQ1DN) steps, respectively.

Step 8 (PG): Bed 1 is now receiving purge gas (PG) from bed 4, and beds 2 and 3 are undergoing the second adsorption step and first product pressurization step (PP1), respectively.

Step 9 (EQ1UP): Bed 1 is undergoing the first equalization rising step (EQ1UP) by receiving low pressure equalization gas from bed 4 that is undergoing the second equalization falling step (EQ2DN). During the same time, beds 2 and 3 are undergoing the third adsorption step (AD3) and the second product pressurization step (PP2), respectively.

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Step 10 (EQ2UP): Bed 1 is undergoing the second equalization rising step (EQ2UP) by receiving high pressure equalization gas from bed 2 that is undergoing the first equalization falling step (EQ1DN). During the same time, beds 3 and 4 are undergoing the first adsorption step (AD1) and countercurrent blowdown step (BD), respectively.

Step 11 (PP1): Bed 1 is receiving the first product pressurization (PP1) gas from bed 3 that is also in the second adsorption step (AD2), while bed 2 is undergoing the cocurrent depressurization step to provide purge gas (PPG) to bed 4.

Step 12 (PP2): Bed 1 is receiving second product pressurization (PP2) gas from bed 3 that is also in the third adsorption step (AD3). During the same time, bed 2 undergoes a second equalization falling step (EQ2DN) by sending low pressure equalization gas to bed 4 that is undergoing the first equalization rising (EQ1UP) step.

A summary of the aforementioned twelve-step PSA process is given in Table I below. In particular, Table I summarizes the valve sequence over one complete cycle for the four-bed PSA system shown in Figure 6. Note from Table 1 that the four beds PSA process operates in parallel, and during $\frac{1}{4}$ of the total cycle time one of the beds is in the adsorption step, while the other beds are either undergoing pressure equalization, purge, blowdown, or product pressurization.

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Table 1: Four-Bed H₂ PSA Valve Switching

(0 = OPENED, C = CLOSED)

Step	1	2	3	4	5	6	7	8	9	10	11	12
Time (Sec)	40	85	25	40	85	25	40	85	25	40	85	25
Bed 1 (BD1)	AD1	AD2	AD3	EQ1 DN	PPG	EQ2	BD	PG	EQ1 UP	EQ2 UP	PP1	PP2
Bed 2 (BD2)	BD	PG	EQ1 UP	EQ2 UP	PPI	PP2	AD1	AD2	AD3	EQ1 DN	PPG	EQ2 DN
Bed 3 (BB3)	EQ1 DN	PPG	EQ2 DN	BD	PG	EQ1 UP	EQ2 UP	PP1	PP2	ADI	AD2	AD3
Bed 4	EQ2 UP	PP1	PP2	AD1	AD2	AD3	EQ1 DN	PPG	DN	BD	PG	EQ1 UP
Valve No.												
31	0	0	0	C	C	C	C	C	C	C	C	C
32	C	C	C	C	C	C	0	0	0	C	C	C
33	C	C	C	C	C	C	C	C	C	0	0	0
34	C	C	C	0	0	0	C	C	C	C	C	C
35	0	0	C	0	0	C	0	0	C	0	0	C
36	C	C	C	C	C	C	0	0	C	C	C	C
37	0	0	C	C	C	C	C	C	C	C	C	C
38	C	C	C	0	0	C	C	C	C	C	C	C
39	C	C	C	C	C	C	C	C	C	0	0	C
40	C	0	0	C	0	0	C	0	0	C	0	0
41	0	0	0	C	C	C	C	C	C	C	C	C
42	C	C	C	C	C	C	0	0	0	C	C	C
43	C	C	C	C	C	C	C	C	C	0	0	0
44	C	C	C	0	0	0	C	C	C	C	C	C
45	C	C	C	C	0	0	C	0	0	C	C	C
46	C	0	0	C	C	C	C	C	C	C	0	0
47	C	0	0	C	0	0	C	C	C	C	C	C
48	C	C	C	C	C	C	C	0	0	C	0	0
49	C	C	C	0	C	C	C	C	C	0	0	0
50	C	C	C	0	0	0	C	C	C	0	C	C
51	0	C	C	C	C	C	0	0	0	C	C	C
52	0	0	0	C	C	C	0	C	C	C	C	C

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It should be noted that the twelve-step PSA cycle is illustrative only, and is given to demonstrate the enhanced PSA process performance achieved by replacing 5A with the CaX adsorbent in the upper layer of the adsorbent beds shown in Figure 6. Other PSA cycles may also be used to show the enhanced PSA process performance obtained in accordance with the invention without deviating from its scope.

A detailed adsorption model, based upon the governing material and energy balances involved in the process, has been applied to simulate the PSA process described above. A plug flow with negligible axial dispersion was used in all of the PSA simulations. Additional characteristics of the model include: bed pressure drop, multicomponent isotherm (as determined by the loading ratio correlation), adiabatic energy balance and adsorption rate (as determined by the linear driving force). The simulation results were found to agree well with pilot scale experimental process performance results.

The results shown in Examples 1-4 below were obtained from PSA simulation results using the following feed mixture on a dry basis: 74.03% H₂, 22.54% CO₂, 0.36% CO, 2.16% CH₄ and 0.91% N₂. Also, total bed size factor is the total quantity of adsorbents per ton per day of 1-12 produced.

Example 1 - Use of CaX (2.0) Adsorbent

Table 2 below discloses the operating conditions and PSA performance for a process utilizing CaX (2.0) adsorbent in the top layer of each of the adsorbent beds B1 to B4 in the system illustrated in Figure 6 of

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the drawings. The process is performed in the manner set forth in Table 1 and shown in Figures 7 and 8 of the drawings. In Tables 2-5, the symbols have the following meaning: TPD = ton per day of hydrogen, s = time unit in seconds, ton 2000 lb.

TABLE 2 - Example 1

Cycle time(s)	600
Adsorbent in first layer of Bed	Alumina
Amount of alumina (lb/TPD H ₂)	1.0385 X 10 ³
Adsorbent in second layer of bed:	activated carbon
Amount of activated carbon (lb/TPD H ₂)	4.9170 X 10 ³
Adsorbent in third layer of bed:	CaX (2.0) zeolite
Amount of CaX (2.0) zeolite (lb/TPD H ₂)	1.5102 X 10 ³
High Pressure: kPa	1.171 X 10 ³
Low Pressure: kPa	1.327 X 10 ²
Feed Flux: ² kmol/s m ²	1.392 X 10 ⁻²
Hydrogen Purity:	99.993%
Hydrogen Recovery:	81.6%
Total Bed Size Factor (lb/TPD H ₂):	7.4657 X 10 ³
Temperature: K	311

Example 2 - Use of LiX Adsorbent

Table 3 below discloses the operating conditions and PSA performance for a process utilizing a LiX zeolite adsorbent in the top layer of each of the adsorbent beds B1 to B4, carrying out the process in the same manner as in Example 1.

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TABLE 3 - Example 2, LiX Zeolite

Cycle time(s)	600
Adsorbent in first layer of Bed	Alumina
Amount of alumina (lb/TPD H ₂):	1.0645 X 10 ³
Adsorbent in second layer of bed:	activated carbon
Amount of activated carbon (lb/TPD H ₂):	5.0400 X 10 ³
Adsorbent in third layer of bed:	LiX zeolite
Amount of LiX zeolite (lb/TPD H ₂):	2.5801 X 10 ³
High Pressure: kPa	1.171 X 10 ³
Low Pressure: kPa	1.327 X 10 ²
Feed Flux: kmol/s m ²	1.392 X 10 ⁻²
Hydrogen Purity:	99.993%
Hydrogen Recovery:	79.61%
Total Bed Size Factor (lb/TPD H ₂):	8.6845 X 10 ³
Temperature:K	311

Example 3 - Use of VSA6 Adsorbent

Table 4 below discloses the operating conditions and PSA performance for a process utilizing a VSA6 zeolite adsorbent in the top layer of each of the adsorbent beds B1 to B4, and again carrying out the process in the same manner as in Example 1.

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TABLE 4 - Example 3, VSA6 Zeolite

Cycle time(s)	600
Adsorbent in first layer of Bed	Alumina
Amount of alumina (lb/TPD H ₂):	1.0568 X 10 ³
Adsorbent in second layer of bed:	activated carbon
Amount of activated carbon (lb/TPD H ₂):	5.0035 X 10 ³
Adsorbent in third layer of bed:	VSA6 zeolite
Amount of VSA6 zeolite (lb/TPD H ₂):	2.3906 X 10 ³
High Pressure: kPa	1.171 X 10 ³
Low Pressure: kPa	1.327 X 10 ²
Feed Flux: kmol/s m ²	1.392 X 10 ⁻²
Hydrogen Purity:	99.984%
Hydrogen Recovery:	80.19%
Total Bed Size Factor (lb/TPD H ₂):	8.4509 X 10 ³
Temperature: K	311

Example 4 Control - Comparative Process Utilizing 5A Zeolite

Table 5 below discloses the operating conditions and PSA performance for a process utilizing a 5A zeolite adsorbent in the top layer of each of the adsorbent beds B1 to B4, carrying out the process in the same manner as in Example 1:

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TABLE 5 - Control 5A Zeolite

Cycle time(s)	640
Adsorbent in first layer of Bed	Alumina
Amount of alumina (lb/TPD H ₂):	1.2108 X 10 ³
Adsorbent in second layer of bed:	activated carbon
Amount of activated carbon (lb/TPD H ₂):	5.7326 X 10 ³
Adsorbent in third layer of bed:	5A zeolite
Amount of 5A zeolite (lb/TPD H ₂):	7.0511 X 10 ³
High Pressure: kPa	1.171 X 10 ³
Low Pressure: kPa	1.327 X 10 ²
Feed Flux: kmol/S.m ²	1.392 X 10 ⁻²
Hydrogen Purity:	99.991%
Hydrogen Recovery:	70.0%
Total Bed Size Factor (lb/TPD H ₂):	1.3995 X 10 ⁴
Temperature: K	311

As shown in Table 6 below and in Figures 9 and 10, use of the CaX (2.0) zeolite adsorbent provides substantially superior results as compared with use of the LiX, VSA6 and 5A zeolites. On the other hand, the use of LiX and VSA6 zeolites results in substantially less adsorbent to provide substantially greater H₂ recoveries and substantially 15 smaller Bed Size Factors than 5A zeolite.

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TABLE 6 - Comparative Performance of CaX (2.0) Zeolite and Other N₂-Selective Zeolite Adsorbents

	Example 1	Example 2	Example 3	Control
N ₂ Adsorbent	CaX (2.0)	LiX	VSA6	5A
Amount of N ₂ Adsorbent (lb./TPD H ₂)	1.5102 x 10 ³	2.5801 x 10 ³	2,3906 x 10 ³	7.0511 X 10 ³
H ₂ Purity	99.993%	99.993%	99.984%	99.991%
H ₂ Recovery	81.6%	79.61%	80.19%	70.0%
Total Bed Size Factor (lb./TPD H ₂)	7.4657 x 10 ³	8.6845 x 10 ³	8.4509 x 10 ³	1.3995 x 10 ⁴

The results of Table 6 and Figure 9 indicate H₂ recoveries of about 70% for 5A zeolite and 80% for the other adsorbents, for about the same H₂ Purity (99.99%). CaX(2.0) allows more feed gas per unit weight of adsorbent at a given P/F ratio (purge to feed) than any of the other adsorbents compared above, i.e. CaX(2.0) has the lowest bed size factor (BSF).

As shown in Figure 10, 78% less zeolite is required as compared to 5A by using CaX (2.0) zeolite, while a 65% reduction is achieved if LiX or VSA6 is used instead of 5A zeolite to produce H₂ at the same purity (99.99%) and recovery (>78%). The smaller bed size factor resulting from the use of CaX reduces the overall void volume such that less hydrogen is lost during the regeneration of the bed, i.e. there is higher H₂ recovery. It is further observed from Figure 8 that N₂ is primarily removed in the CaX zeolite layer (Nodes 32-49) of the bed, while the other impurities (e.g., CO₂, CH₄ and CO) are removed in the alumina (nodes 1- 17) and activated carbon (nodes 17-32) layers upstream of the zeolite layer.

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The superior adsorption characteristics of CaX (2.0) zeolite is further illustrated in Figure 11 of the drawings, which shows the differential loading or working capacity of N₂ on CaX (2.0) as compared with the 5A, LiX and VSA6 zeolites. The differential N₂ loading on CaX (2.0) is more than 5 times, and the differential N₂ loading on LiX and VSA6 more than 3 times that on the 5A zeolite. The differential N₂ loading on each adsorbent in Figure 11 was calculated using the same conditions given above for Figure 3. As shown in the comparative tests and the foregoing Figures, in accordance with the present invention, CaX (2.0) is the adsorbent of choice for the H₂ PSA process of the invention.

Although the foregoing PSA processes relate to H₂ production, the key features of this invention may be extended to other separation processes, e.g. CO₂ production from synthesis gas or other sources containing CO₂ in the feed, or in other PSA processes for the co-production of H₂ and CO. For example, CaX (2.0) could replace 5A or LiX zeolite in other separation processes that require the removal of trace or low concentration levels of N₂ to achieve enhanced PSA process performance and high purity product.

In addition, the zeolite layer/zone of each adsorbent bed may be replaced with multiple layers of different adsorbents. For example, the homogeneous zeolite layer may be substituted by a composite adsorbent layer containing different adsorbent materials positioned in separate zones and employing temperature conditions favoring adsorption performance

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of the particular adsorbent materials under applicable processing conditions in each zone.

It will be understood that these and other changes may be made in the preferred parameters of the PSA process hereof without departing from the invention. Accordingly, it is intended that the scope of this invention should be determined from the claims appended hereto.

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The embodiments of the invention in which an exclusive property or privilege is claimed are defined as follows:

1. A pressure swing adsorption (PSA) process for purifying a feed stream containing more than 50 mole % hydrogen, which comprises passing the feed stream at a pressure above atmospheric pressure through a multilayer adsorbent bed, wherein at least one contaminant from the group H₂O, CO₂, CH₄ and CO is adsorbed from the gas stream prior to passing the stream through a layer of naturally occurring zeolite selected from the group consisting of chabazite, erionite, clinoptilolite and faujasite zeolite, or a layer of synthetic zeolite adsorbent having a SiO₂/Al₂O₃ ratio within the range of 2.0-2.5 and selected from the group consisting of CaX, LiA, LiX and VSA6 adsorbent to adsorb substantially all of the nitrogen in such stream and recovering purified (>99.9%) hydrogen as product from the multilayer adsorbent bed.

2. The pressure swing adsorption process of claim 1, wherein the feed stream contains less than 3% N₂ and preferably less than 1.5% N₂

3. The pressure swing adsorption process of claim 1, wherein the, gas stream immediately prior to being passed through the zeolite adsorbent contains on an average less than 0.15 mole % CO₂.

4. The pressure swing adsorption process of claim 1, wherein the feed gas stream to be treated is a synthesis gas containing from 60 to 90 mole % hydrogen.

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5. The pressure swing adsorption process of claim 1, wherein the zeolite is type X with ion exchange > 90% Ca.

6. The pressure swing adsorption process of claim 1, wherein the feed gas stream is first passed through an adsorbent bed containing an alumina layer for adsorption of H₂O, then through an activated carbon layer for adsorption of CO, CH₄ and CO₂ and then through a zeolite layer for adsorption of nitrogen.

7. The pressure swing adsorption process of claim 1, wherein H₂O, CO₂, CH₄ and CO are substantially adsorbed from the gas stream prior to the stream passing through the naturally occurring adsorbent or the synthetic zeolite adsorbent.

8. The pressure swing adsorption process of claim 1, which comprises passing said stream at pressures of from 5 to 20 bars through four adsorbent beds each of which includes a layer of said naturally occurring or synthetic zeolite for adsorbing nitrogen from the gas stream, utilizing the 12-step PSA step sequence illustrated in Table 1 thereof.

9. The pressure swing adsorption process of claim 1, wherein the total bed size factor is less than 9,000 lb/TPD of hydrogen, and hydrogen recoveries of the order to 80% or greater are obtained.

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10. A PSA system for purifying a feed gas stream; wherein each adsorbent bed comprises:

- a. an adsorbent layer for removing H₂O;
- b. an adsorbent layer for removing carbon dioxide; and
- c. an adsorbent layer of CaX, LiA, LiX or VSA6 zeolite adsorbent having a SiO₂/Al₂O₃ ratio within the range of 2.0-2.5 for N₂ removal.

REFORMING OF NATURAL GAS

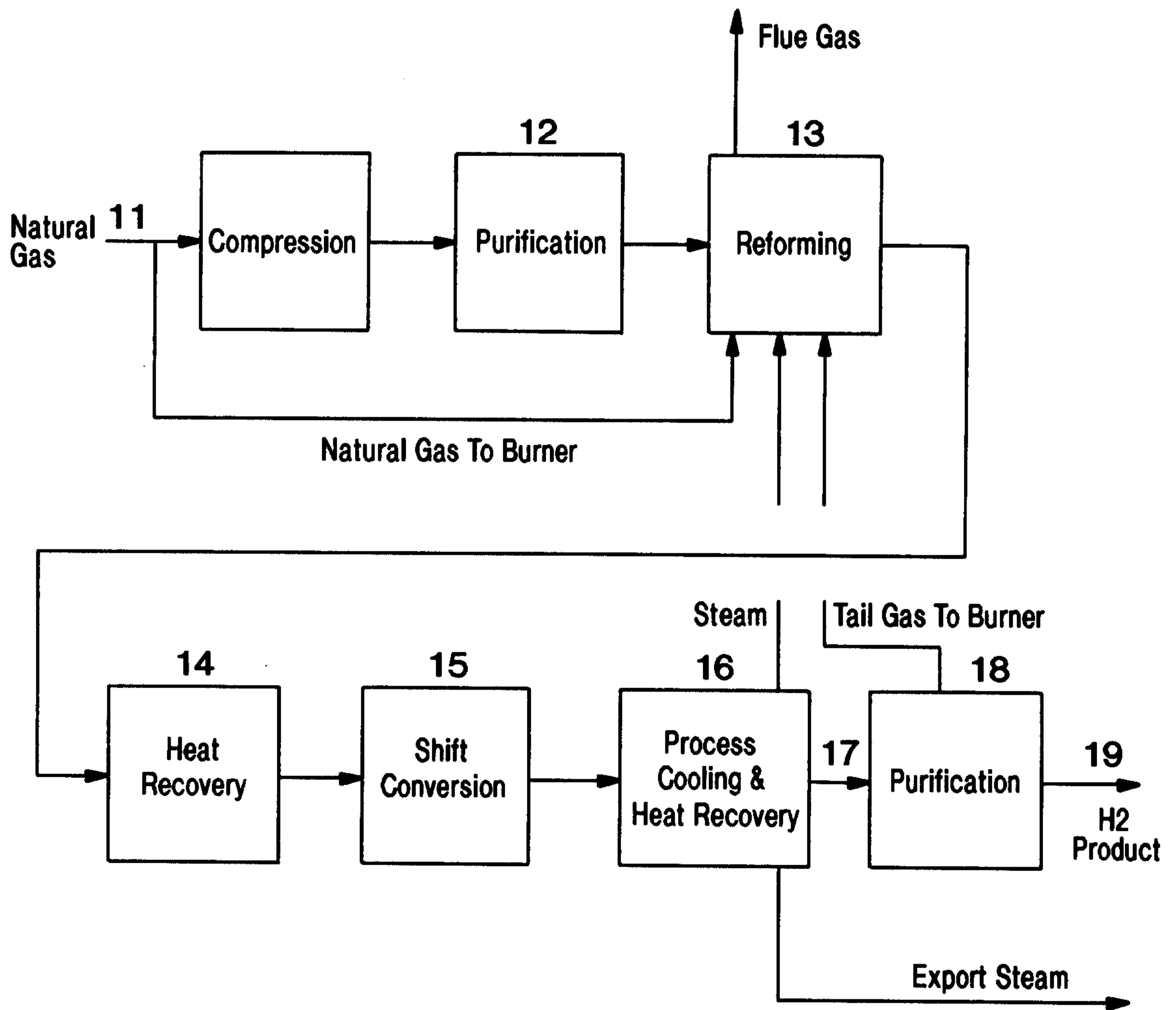


FIG. 1
PRIOR ART

Comparison of the Nitrogen Adsorption Isotherms
on CaX(2.0), Li-X, VSA6 and 5A Zeolites

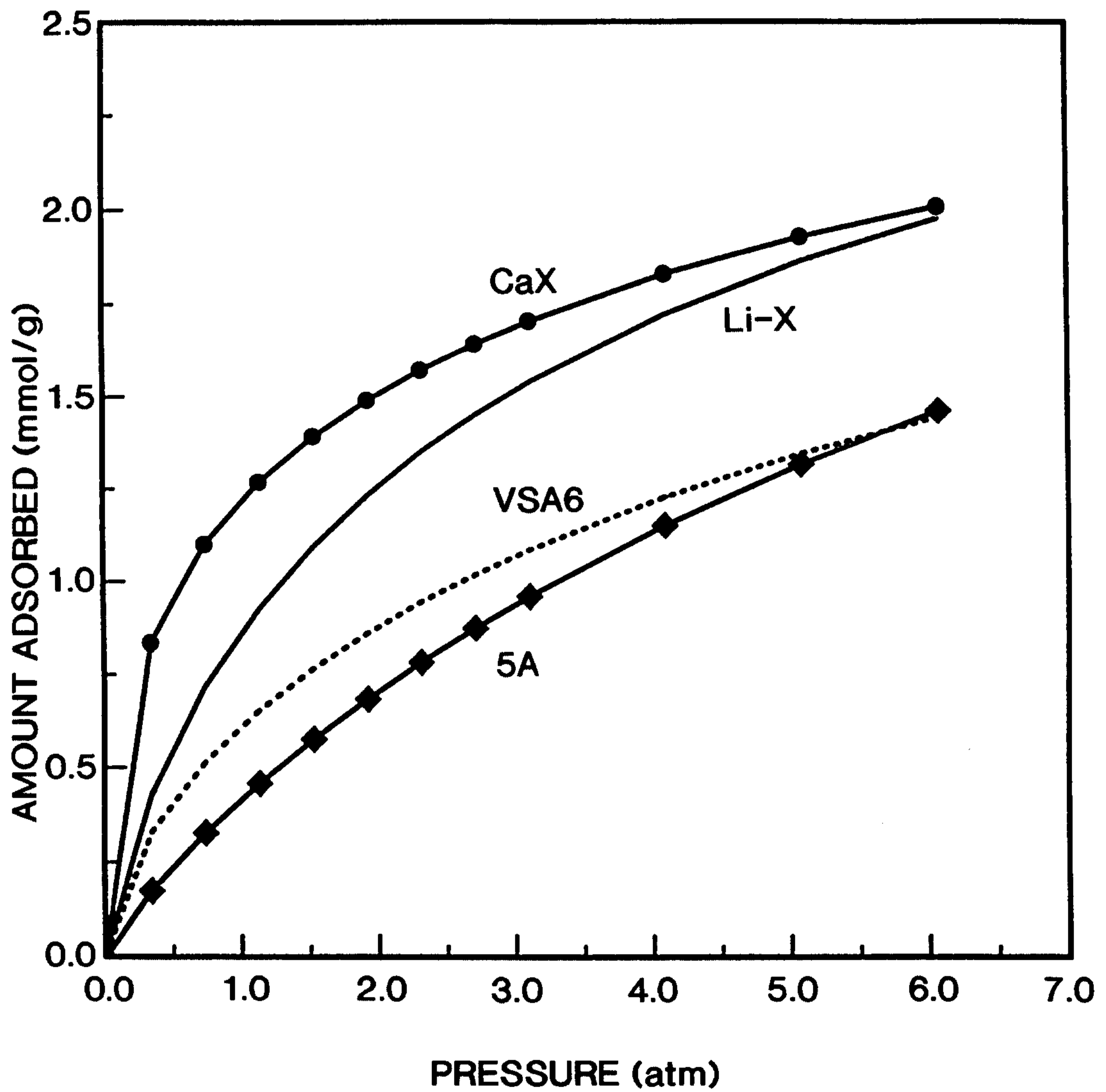


FIG. 2

Comparison of the Differential N₂ Loading of
CaX(2.0) With Other Adsorbents

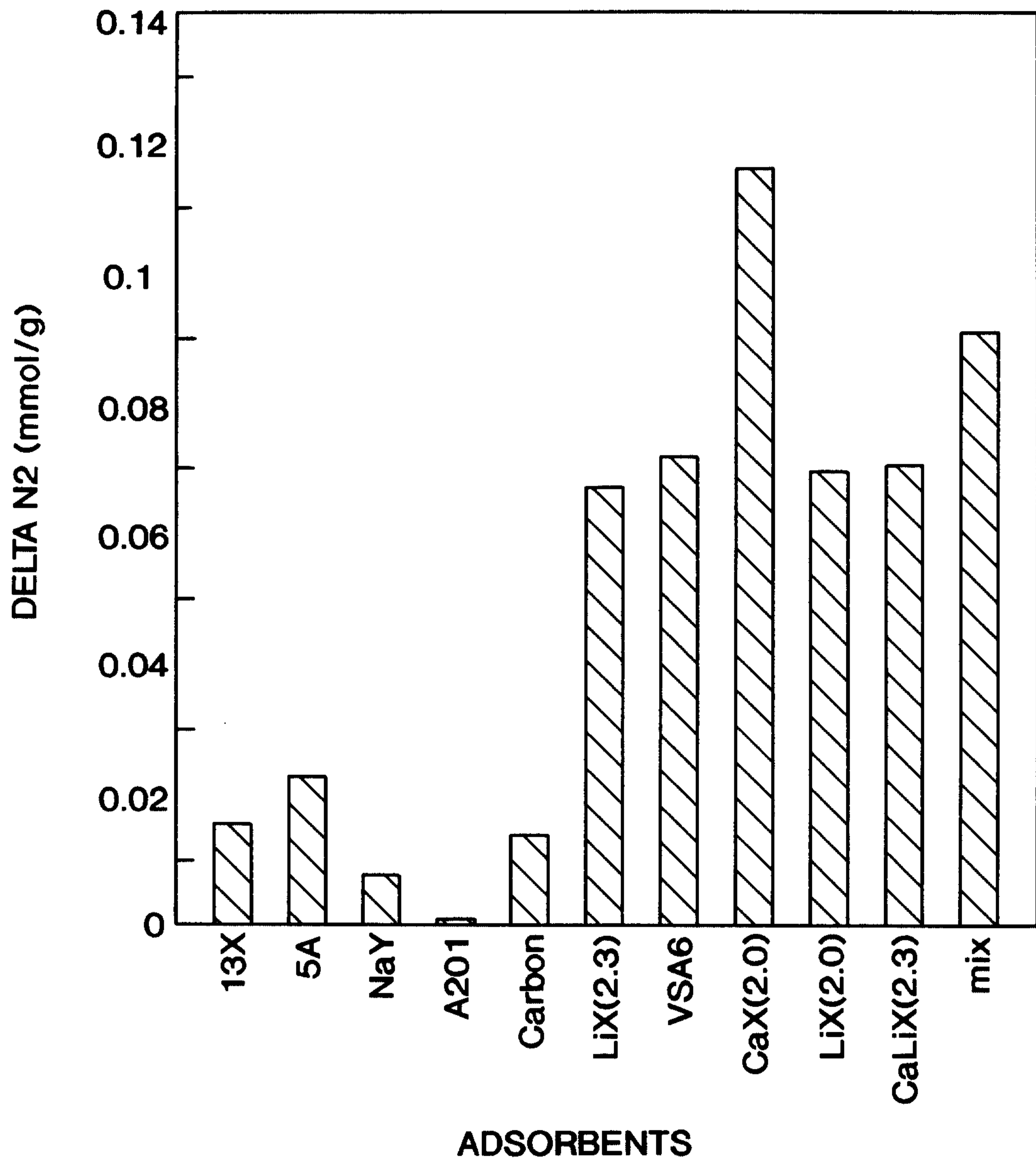


FIG. 3

Comparison of CO₂ Adsorption Isotherms on CaX(2.0) and Other N₂-selective Adsorbents at 300K.

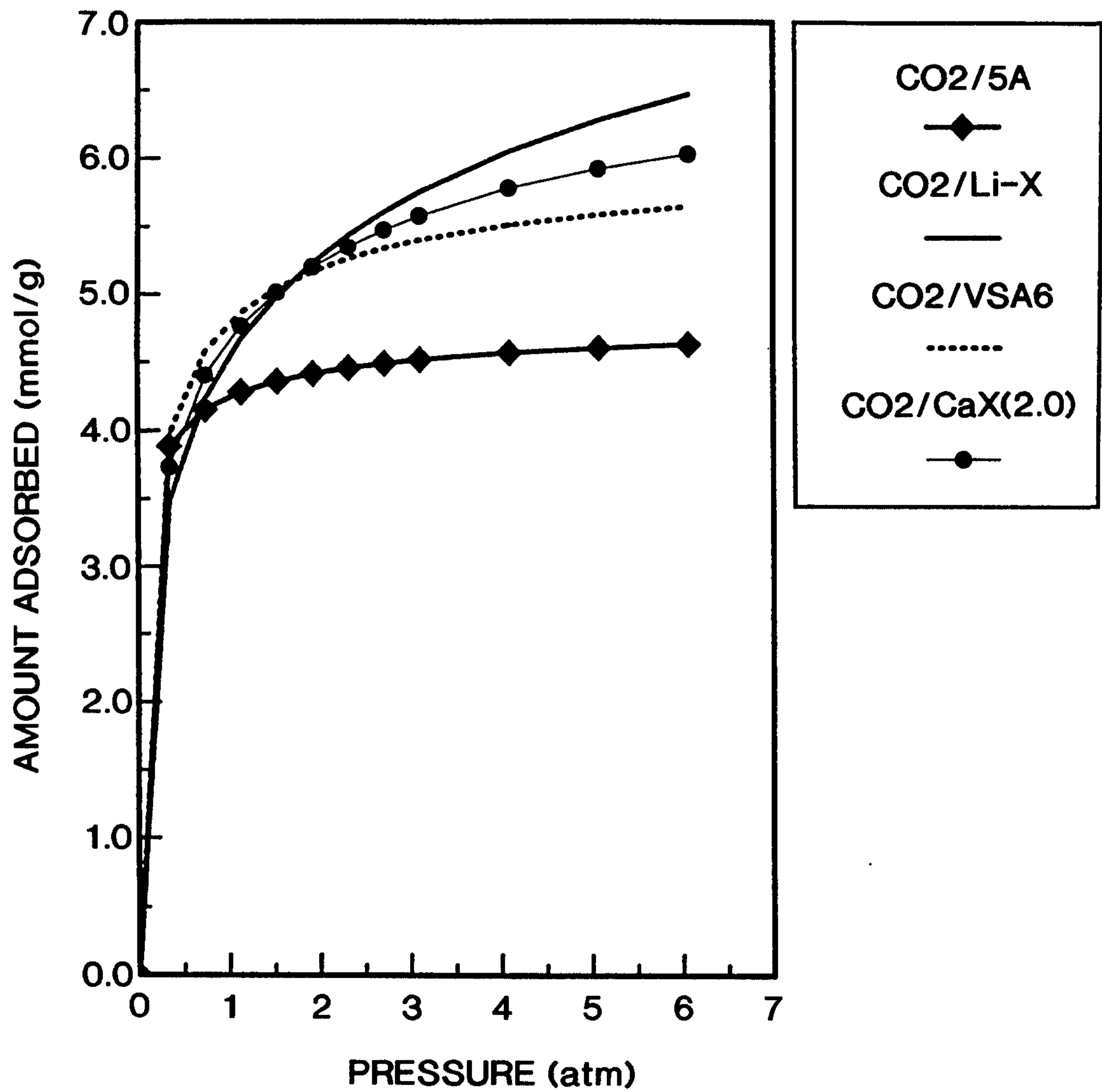


FIG. 4

Schematic Drawing of a PSA Adsorption Bed of the Present Invention.

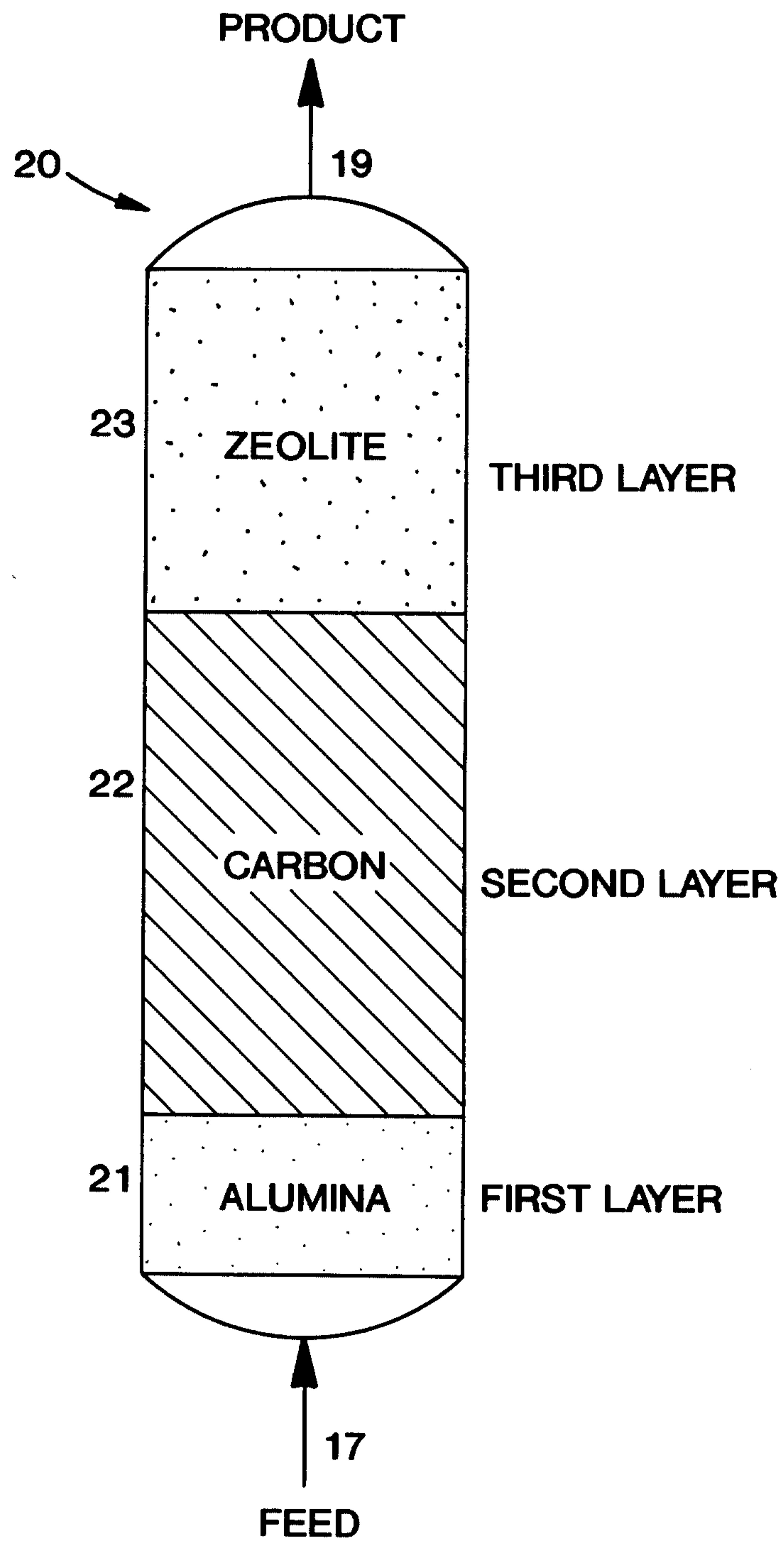


FIG. 5

Schematic Drawing of a Four-Bed System for Carrying out the PSA Process of the Invention.

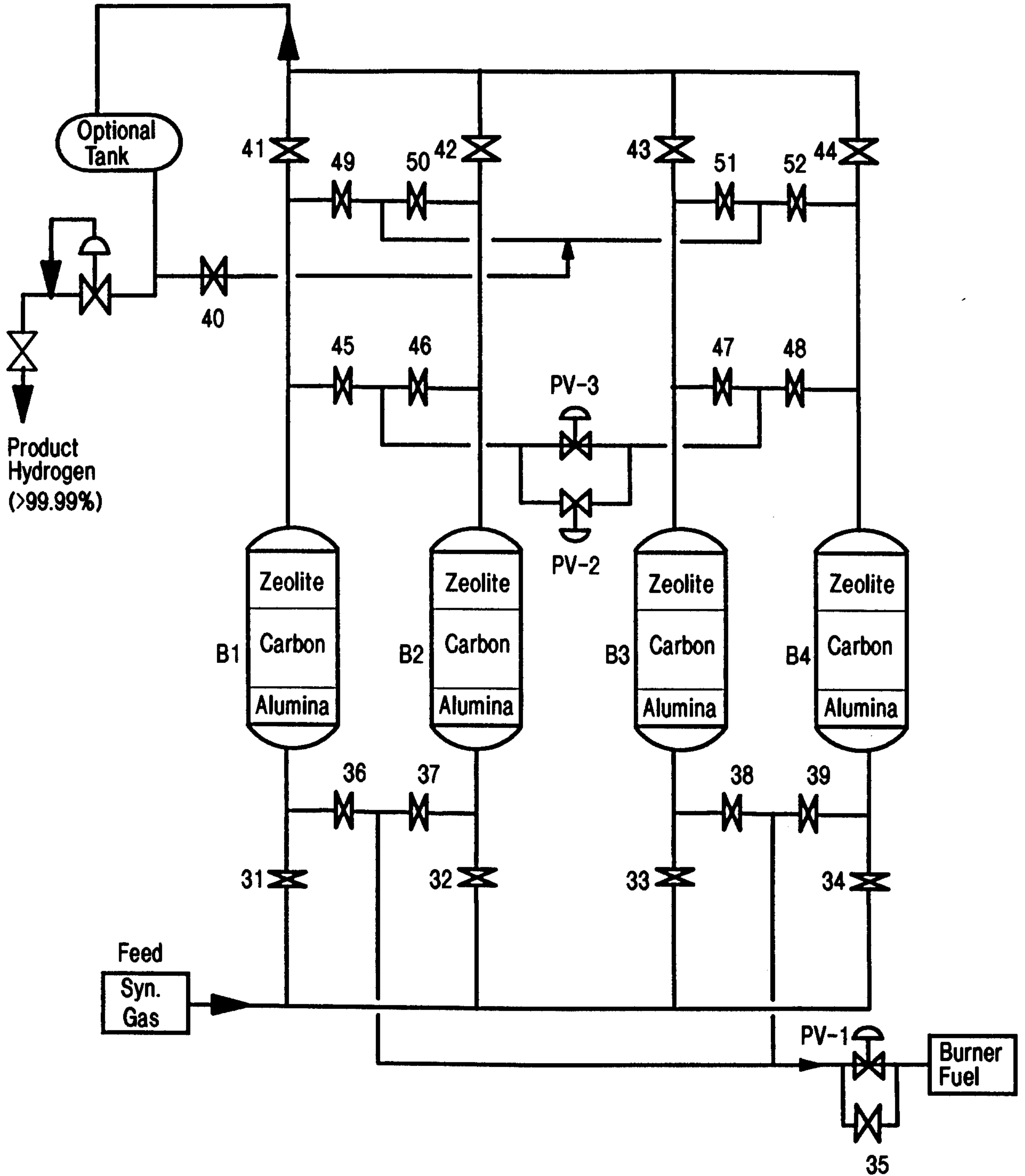


FIG. 6

A Graphic Depiction of the Representative Bed Pressure Profile During One Complete PSA Cycle in the Preferred Embodiment of the Invention Described Herein.

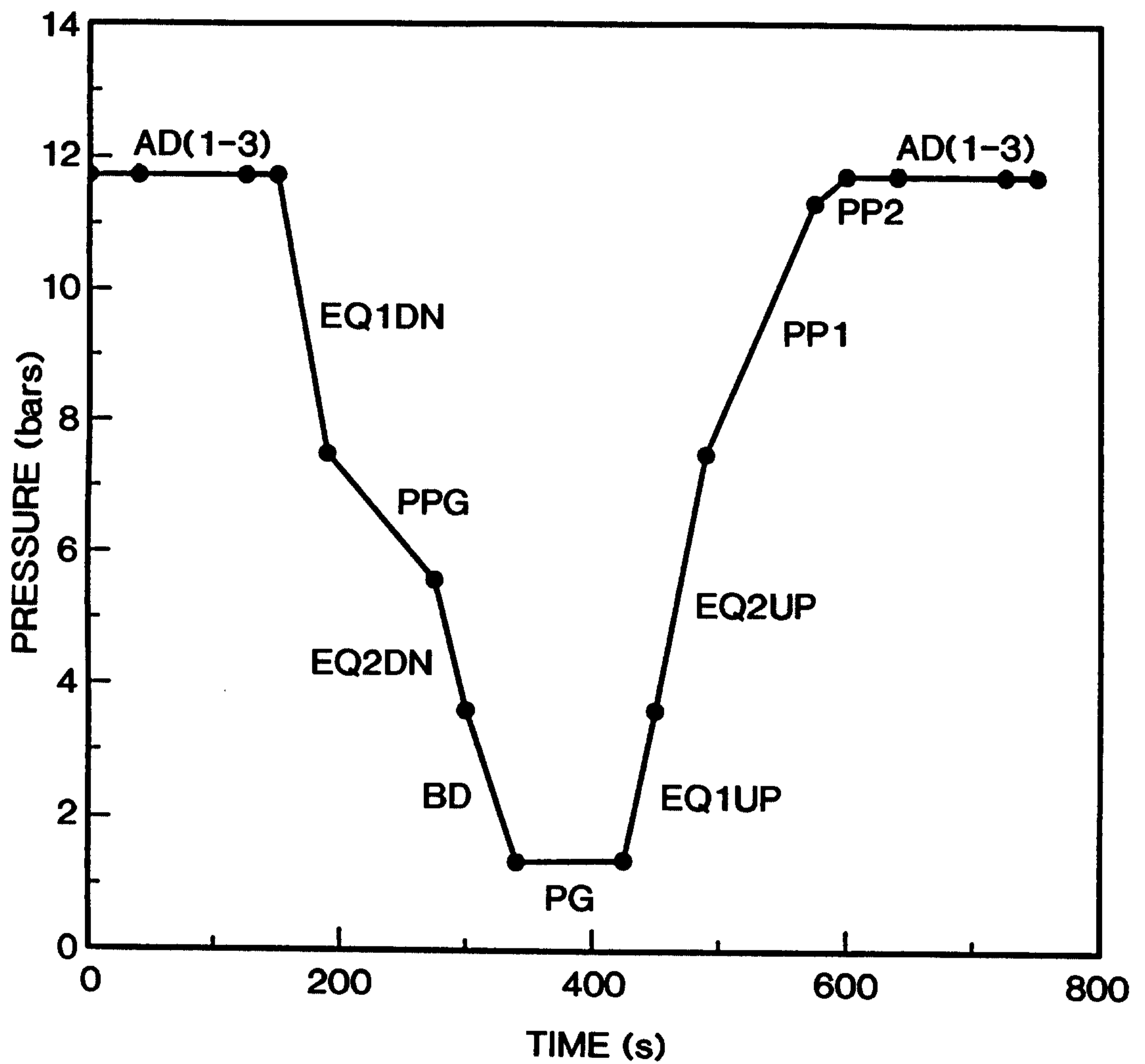


FIG. 7

A Graphic Illustration of the Representative Gaseous Component Mole Fractions in an Adsorbent Bed in the Embodiment Described, at the Conclusion of the High Pressure Adsorption Step Therein.

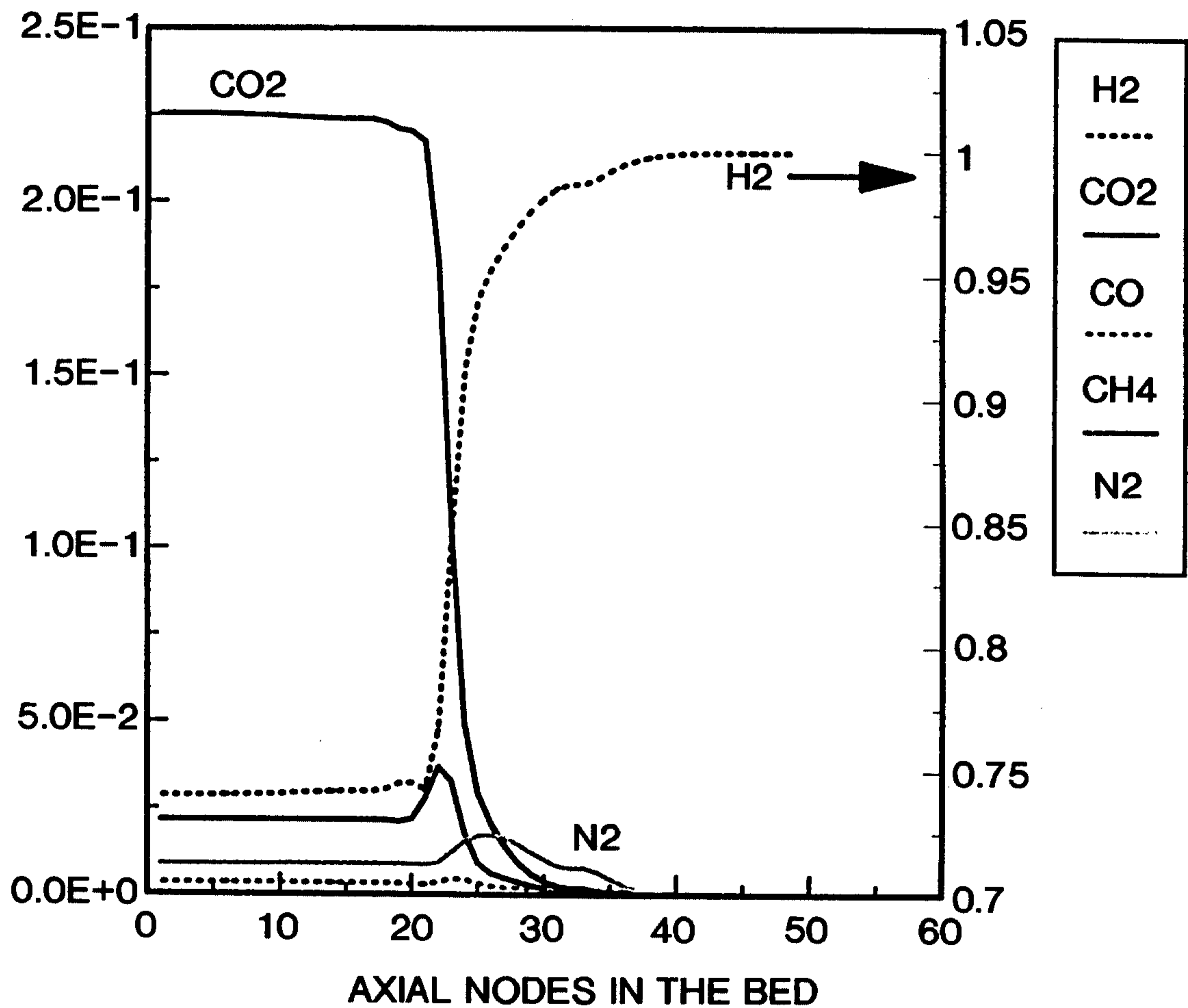


FIG. 8

A Graphic Comparison of the Computer Simulated PSA Process Performance for the System of Figure 4, Utilizing each of the CaX(2.0), 5A, LiX and VSA6 Zeolite Adsorbents in the N₂-Selective Layer.

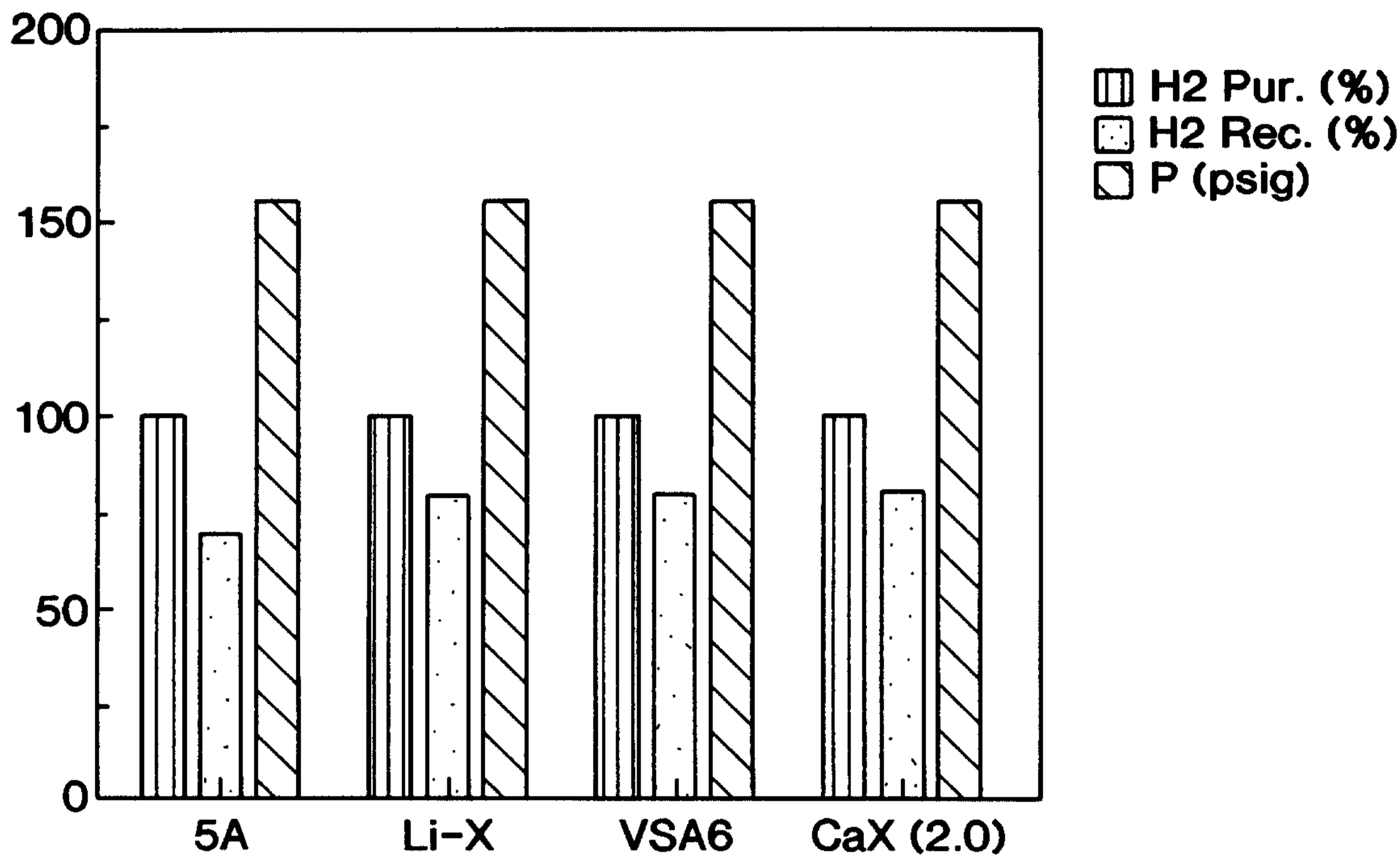


FIG. 9A

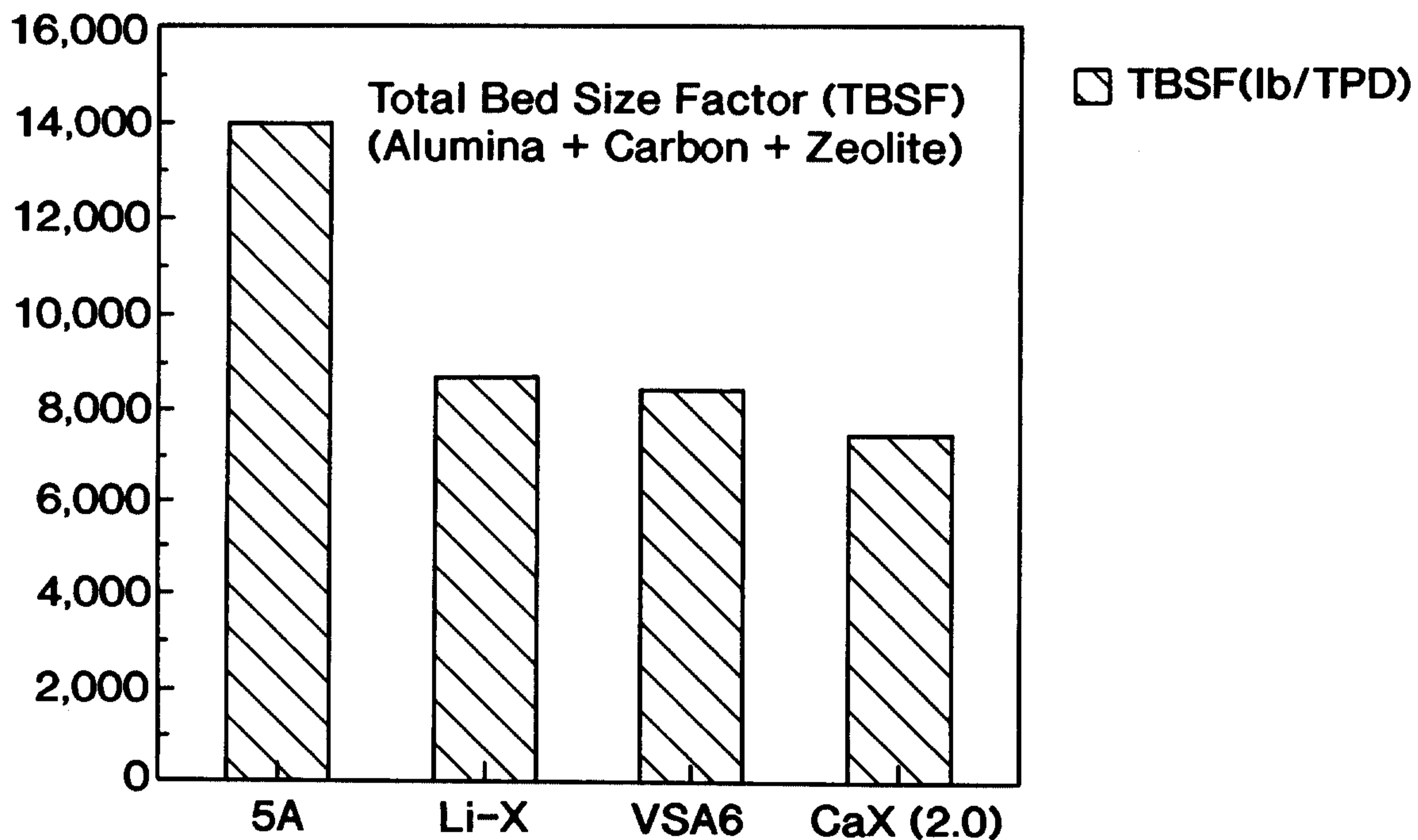


FIG. 9B

A Graphic Comparison of the Quantity of CaX(2.0), 5A, LiX and VSA6 Zeolite Adsorbents in the Illustrated Four Bed PSA Process.

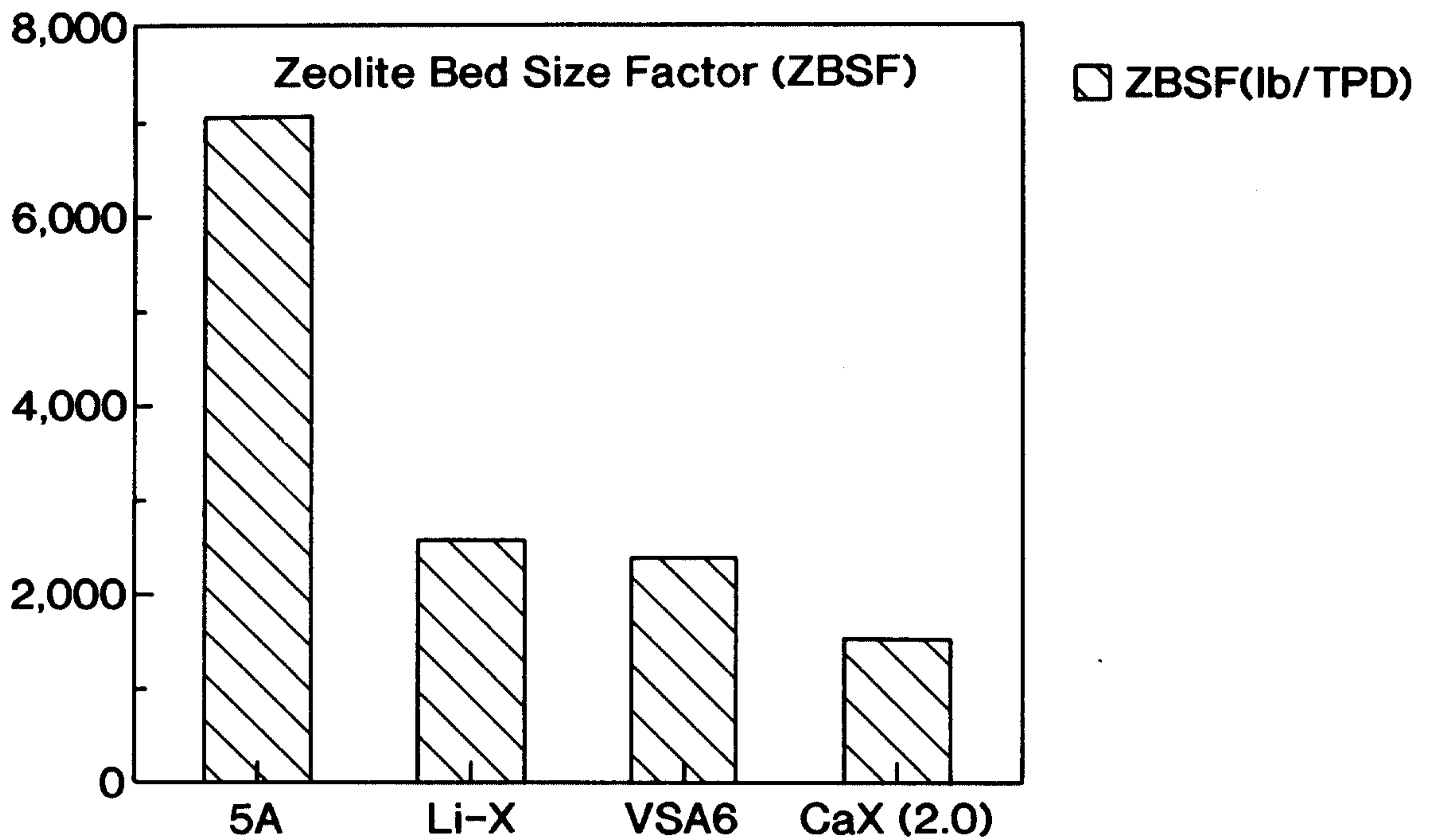


FIG. 10A

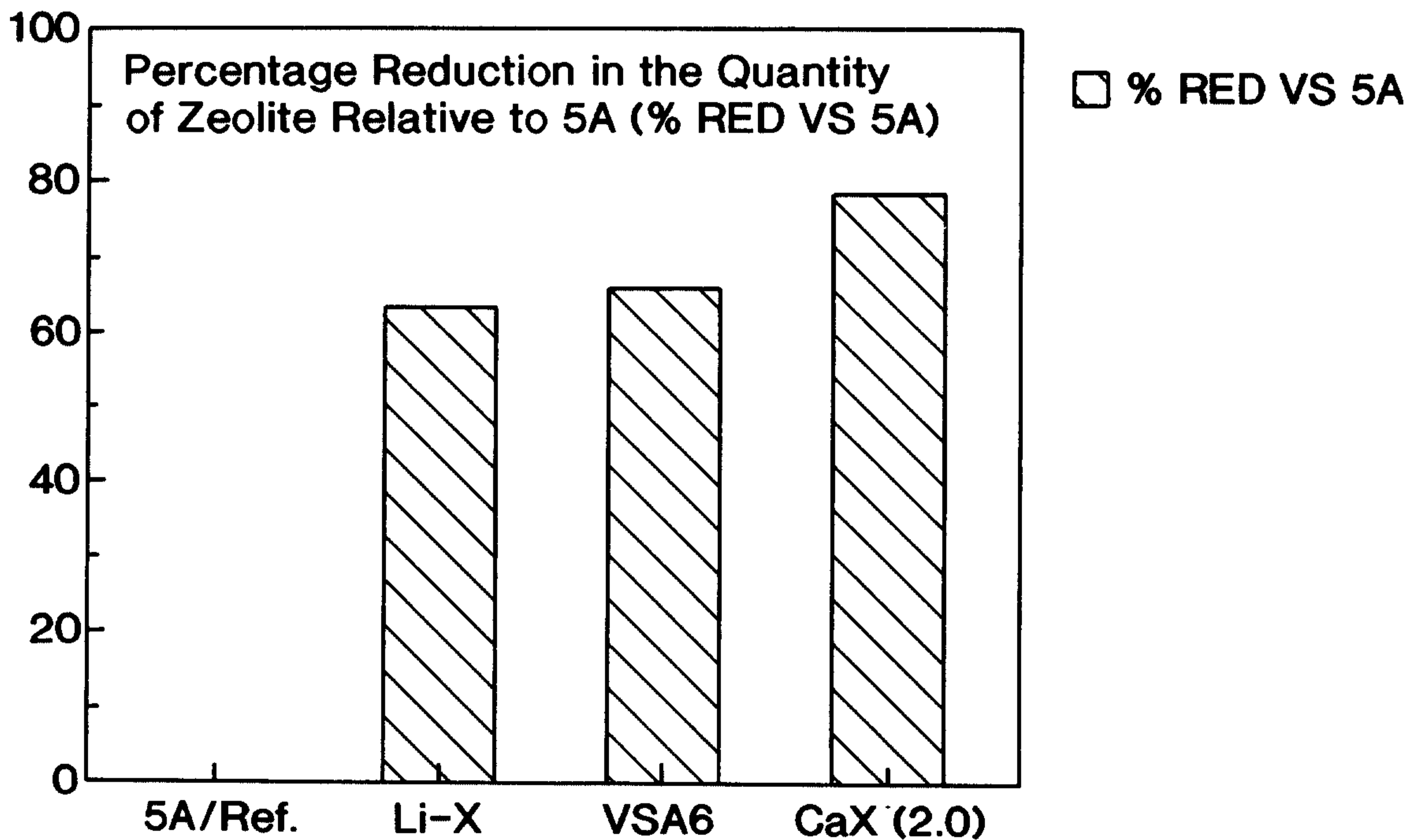


FIG. 10B

A Graphic Comparison of Differential Loading of N₂ on CaX(2.0), 5A, LiX and VSA6 Zeolites.

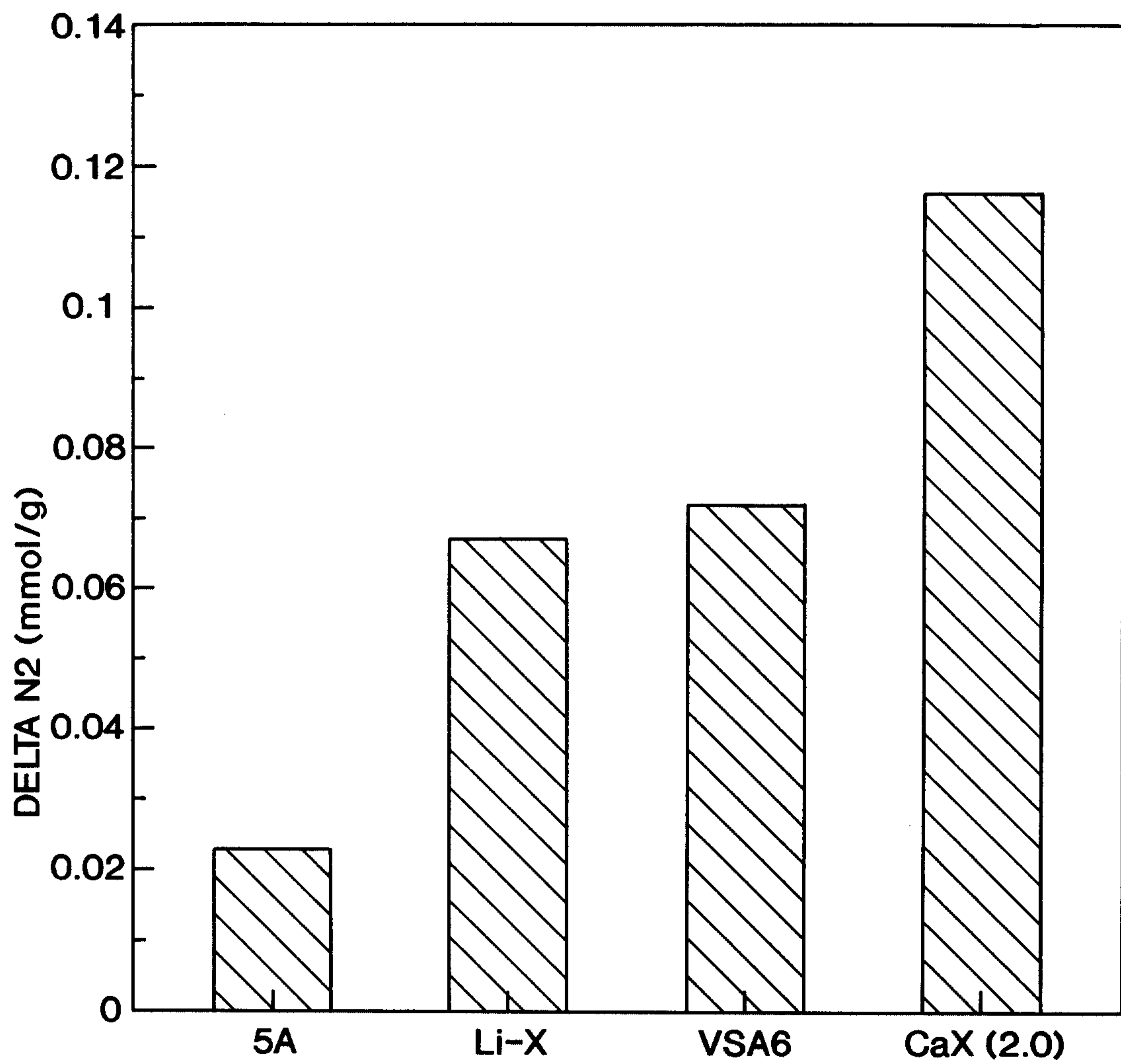


FIG. 11

