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# Hydroquinone clathrate based gas separation (HCBGS): Application to the $CO_2/CH_4$ gas mixture

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# ABSTRACT

Hydroquinone (HQ) clathrates have recently been identified as promising candidates for selective gas capture and storage processes. This study evaluates the effectiveness of HQ clathrates in the separation of  $CO_2$  from  $CO_2/$ CH<sub>4</sub> gas mixtures, through direct gas/solid reactions in a fixed-bed reactor. The influence of the process operating parameters (i.e. reaction time, pressure, temperature and feed gas composition) on the  $CO_2$  capture kinetics, selectivity towards  $CO_2$ , and transient storage capacity were investigated. The experiments were performed using either pure HQ or HQ-based composite materials, with temperatures ranging from about 283 to 343 K, pressures from 3.0 to 9.0 MPa, and  $CO_2$  mole fraction in the gas mixture ranging from 0.2 to 1. The experimental results show that over the range of gas composition investigated, the enclathration reaction is selective to  $CO_2$ . This preferential  $CO_2$  capture is enhanced at high  $CO_2$  mole fractions, low temperatures and high pressures. Regarding gas capture kinetics, it was confirmed that the composite material is much more efficient than pure HQ crystals. The  $CO_2$  enclathration rate increases with temperature, pressure and  $CO_2$ fraction in the feed gas. For the first time, the feasibility of such gas separation techniques using HQ clathrates was demonstrated at bench scale.

#### 1. Introduction

Keywords:

Clathrate

Separation

CO<sub>2</sub> capture

Hydroquinone

Natural gas treatment

Natural gas is the 3<sup>rd</sup> energy source after coal and oil, according to the International Energy Agency [1]. It is mainly composed of CH<sub>4</sub>, but it can also contain other compounds including low molecular weight hydrocarbons (such as ethane, propane, butane, pentane), acid gases (such as CO2 and H2S), incondensable gases (N2 and O2), and traces of rare gases, mercury and water [2]. Light hydrocarbons (especially CH<sub>4</sub>) are the main valuable compounds in natural gases. The other con stituents are defined as impurities because of the complications they induce, such as decreasing the gas calorific value, and their operating constraints (e.g. condensation, corrosion and plugging). To respect the required specifications for valorization and commercialization of the natural gas, the removal of these impurities is of paramount im portance. As acid gases are generally the most constraining (due to their toxic and corrosive properties) and most abundant, in terms of im purities, a gas sweetening treatment (i.e. removal of CO<sub>2</sub> and H<sub>2</sub>S) is required. CO<sub>2</sub> in particular one of main gases contributing to green house effects [3] must be captured and stored (e.g. by injection into an underground reservoir) to curb its concentration from increasing in the atmosphere [4]. Actually, several processes are available for re moving CO<sub>2</sub> from natural gas (e.g. chemical absorption using amines,

adsorption on porous materials, permeation through membranes, con densation, and cryogenic separation) [4,5] for which the CO<sub>2</sub> recovery rate is high, reaching values > 80%. However, these processes are limited by economic criteria related to energy consumption [5,6]. In this way, alternative techniques for CO<sub>2</sub> gas capture clathrate based technologies being one of them are subjects of great interest.

Gas clathrates are solid state host guest compounds consisting of a network of self associating molecules forming cavities (called "hosts") in which gas species (called "guests") can be encaged [7,8]. The com bination of host and guest molecules to form clathrates is only possible in appropriate thermodynamic conditions. These inclusion compounds are known to store large amounts of gases such as CO<sub>2</sub>, H<sub>2</sub>S, CO, SO<sub>2</sub>, CH<sub>4</sub> [9], and to be potentially selective to a specific gas present in a mixture [8]. This latter property has motivated many researchers to investigate the potential of clathrates in gas separation. The principle of this concept is based on the simple fact that when a gas clathrate is formed from a gas mixture, one of the constituents (e.g.  $CO_2$  in a  $CO_2/$ CH<sub>4</sub> mixture) may be selectively encaged in the solid structure. A clathrate based separation process would thus comprise a clathrate formation step followed by a dissociation step: (i) the formation step leads to a residual vapor and a solid clathrate under equilibrium, both having different gas compositions (e.g. the gas containing a lower CO<sub>2</sub>

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concentration than the initial gas mixture, and the clathrate a higher one); (ii) the dissociation step: by changing only the thermodynamic conditions (e.g. an increase of temperature, a decrease of pressure, or both), the gaseous molecules contained in the clathrate are released (e.g. producing a  $CO_2$  rich gas mixture). Theoretically, this concept is very attractive, especially when the gas to be treated is already pres surized, such as for production gas. Actually, additional work is truly necessary to demonstrate the feasibility of such separations at larger scale (e.g. using bench and pilot scale apparatuses).

Among the clathrate based processes, the most studied involves gas clathrate hydrates (often simply called gas hydrates), where the host molecules are water molecules associating by hydrogen bonds, gen erally formed at high pressure (several MPa) and moderately low temperature (close to 273 K) [10]. Accordingly, the hydrate based gas separation (HBGS) process has already attracted widespread interest [11 20]. Nevertheless, although the HBGS process may be economic ally competitive (as the enclathration precursor is water, and the pro cess can work under pressure), the deployment of the proof of concept to industrial scale is still limited by numerous critical disadvantages: (i) the additional cost of maintaining a relatively low temperature, re quired in the reactor to form the hydrates; (ii) the slow crystallization rate; (iii) the insufficient capture selectivity, particularly for the CO<sub>2</sub>/ CH<sub>4</sub> gas mixture; (iv) the necessity to add chemical promoters to overcome the thermodynamic and kinetic limitations [19], some of which can be toxic (e.g. tetrahydrofuran) or cause foaming problems during hydrate dissociation (e.g. sodium dodecyl sulfate); and (v) the operating and technological constraints related to the handling and transport of the multiphase mixture under pressure (i.e. slurry com posed by solid hydrates, liquid water, and gases). There are plenty of examples in literature of studies carried out to address these challenges, many of them based on semiclathrates (e.g. TBAB, TBAF, etc.). The main advantage of using these substances is that the HGBS process can be operated at near ambient pressure and/or temperature conditions [20].

Alternatively, hydroquinone (a HQ) is an organic compound from the phenol family, well known to form gas clathrates. Although organic clathrates were discovered over a century ago [21], they were mainly fascinating scientific curiosities, but very few practical applications were proposed to date. A very limited number of studies involving or ganic clathrates is available in literature, in comparison to common gas hydrates. For example, the precise molecular structure of the simple CO<sub>2</sub> HQ clathrate was reported only recently by Torré et al. [22] showing that much remains to be done even at the fundamental level to increase our understanding of such compounds. The stoichiometry of the HQ clathrate ( $\beta$  HQ) is 3:1 (i.e. 3 molecules of HQ per guest mo lecule), leading to a maximum storage capacity of  $3.03 \text{ mol}^{\text{Guest}}/\text{kg}^{\text{HQ}}$ [23]. Recently, Coupan et al. showed that HQ clathrates can be formed with CO<sub>2</sub>, CH<sub>4</sub> and CO<sub>2</sub>/CH<sub>4</sub> mixtures over a wide range of tempera tures (up to about 354 K) and at moderate pressure (< 1 MPa) [24,25]. Moreover, HQ clathrates can form by direct gas solid reaction (i.e. a solvent is not required) by simply contacting solid HQ with the gas in appropriate pressure and temperature conditions [26 28]. As the ki netics of this solid solid transition (i.e. from  $\alpha$  to  $\beta$  HQ) is rather slow, an innovative HQ based composite material had already been devel oped by the authors to primarily improve the enclathration kinetics, as well as to also overcome some process limitations. With the composite: (i) a gas solid enclathration reaction is achieved, for which the induc tion time (i.e. the waiting period before the  $\alpha$  HQ starts to transform into  $\alpha$  HQ) is cancelled and the global kinetics is significantly im proved; (ii) it permits both enclathration formation and additional adsorption phenomena; (iii) it is compatible with industrial gas solid contactors such as fixed bed reactors; (iv) this composite can be re used over several capture/regeneration cycles; and (v) it avoids the handling of HQ powder, the fine particles to be blown in the process unit, and high pressure drop in the gas/solid contactors [27]. Interestingly, var ious authors have reported that the HQ clathrates could be used for the

selective capture of CO<sub>2</sub> from various CO<sub>2</sub> containing gas mixtures (such as CO<sub>2</sub>/H<sub>2</sub>, CO<sub>2</sub>/N<sub>2</sub> and CO<sub>2</sub>/CH<sub>4</sub> gas mixtures) [29 33]. Parti cularly, Coupan et al. and Lee et al. gave clear evidence on the cap ability of HQ clathrates in the selective trapping of CO<sub>2</sub> from an equi molar CO<sub>2</sub>/CH<sub>4</sub> mixture [32 33]. However, only one quantitative selectivity of 29 mol<sup>CO2</sup>/mol<sup>CH4</sup> from the equimolar CO<sub>2</sub>/CH<sub>4</sub> mixture has been reported in literature to date, without any precision on the pressure and temperature condition associated to this result [33].

The present work aims at providing a technical performance eva luation of a HQ clathrate based gas separation (denoted as HCBGS in the following) process applied to a  $CO_2/CH_4$  gas mixture. The  $CO_2/CH_4$ gas mixture and the experimental conditions of this study were chosen to approach the industrial case study of the gas sweetening applications [2,34]. The experiments were performed with CO<sub>2</sub>/CH<sub>4</sub> gas mixtures with CO<sub>2</sub> mole fraction from 0.2 to 1 in the  $\sim$  283 343 K and pressure from 3.0 to 9.0 MPa. The CO<sub>2</sub> capture selectivity, the transient gas storage capacity (i.e. the quantity of gas captured at a given time), and the capture kinetics were assessed. The first part of the study was de voted to the use of pure native HQ. Although we believe that the direct use of such crystals is not suitable for industrial purposes, this config uration represents a case study in which the gas capture is due only to enclathration. Therefore, this medium was used to investigate and un derstand the influence of the process parameters on the separation ef ficiency. In the second part, an HQ based composite was used for the separation. This conditioning has shown to give better results in terms of kinetics than native HQ crystals, but the phenomena involved in the CO<sub>2</sub> capture are more complex as they involve both adsorption (on the porous medium) and enclathration (with HQ). With this composite, the influence of the process parameters was also analyzed and compared against the native HQ crystals.

### 2. Experimental section

#### 2.1. materials

HQ (purity of 99.5 mol%) was purchased from Acros Organics. Helium, CO<sub>2</sub> and CO<sub>2</sub>/CH<sub>4</sub> gas mixtures (minimum mole fraction purity of 99.995%) were purchased from Linde Gas SA. The CO<sub>2</sub> mole frac tions of the CO<sub>2</sub>/CH<sub>4</sub> mixtures were measured by chromatography at 0.204, 0.399, 0.501, 0.601, 0.733 and 0.795. These values are in agreement with those given by the supplier specified within a relative uncertainty of  $\pm$  2%. The mixtures will be referenced in the following by the nominal values, as 0.20, 0.40, 0.50, 0.60, 0.75 and 0.80.

For the experiments, either native HQ or HQ based composite were used. Native HQ was put in contact with pure  $CO_2$  gas at 3.0 MPa and ambient temperature for 1 month to form  $CO_2$  HQ clathrates, followed by a dissociation step at 1 kPa and 343 K until all the stored gas was released. This process generates a preformed form of HQ with a highly reduced induction time [32]. The structure of the preformed HQ was investigated by Raman spectroscopy just before contacting the  $CO_2/$ CH<sub>4</sub> gas mixture (by taking a sample inside the reactor). The analysis revealed the presence of a mixture of  $\alpha$  HQ and guest free  $\beta$  HQ, with the  $\alpha$  HQ structure in majority (the two HQ forms cannot be dosed precisely with such Raman analysis). Moreover, several other Raman spectroscopic analyses achieved on the cross section of preformed HQ crystals revealed that (in these conditions) the crystals present a core shell structure: the guest free  $\beta$  HQ structure forms the external shell while the crystal core is composed of  $\alpha$  HQ.

The HQ based composite used in this work is the one synthesized by Coupan et al. with a dry impregnation process [27]. It consists of Si liasphere<sup>®</sup> silica particles (size 200 500  $\mu$ m, pore size of 100 nm, spe cific area of 57 m<sup>2</sup>/g, and porous volume of 0.83 cm<sup>3</sup>/g) impregnated with HQ. The HQ content is 0.44 g<sup>HQ</sup>/g<sup>Silica</sup>. The composite material was also pre formed with the pure CO<sub>2</sub> in the same conditions as those used for preforming native HQ. As the HQ crystallites are mainly de posited inside the pores of the silica particles, the internal structure of

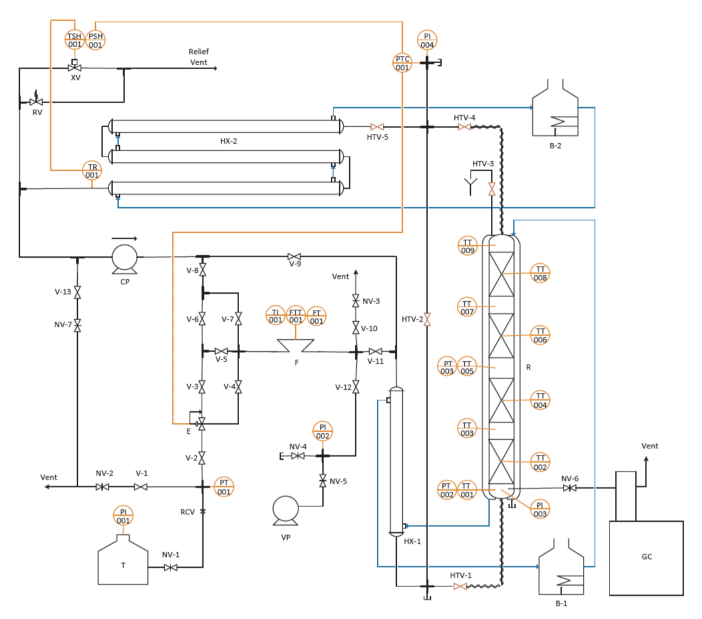


Fig. 1. HCBGS process: (T) gas tank; (E) regulator expander; (CP) gas circulator pump; (VP) vacuum pump; (F) Coriolis flowmeter; (HX) heat exchanger; (R) fixed-bed reactor; (B) thermostatic bath; (GC) gas chromatograph; (V) plug valve; (NV) needle valve; (RCV) rapid connection valve; (HTV) high-temperature plug valve; (RV) relief valve; (XV) pneumatic valve. Blue line refers to heat transfer fluid. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

such nanocrystals cannot be investigated as it was not possible to isolate and cut a crystallite with a scalpel to analyze its cross section). Therefore, following our conclusions obtained with native HQ crystals, we assume that the HQ crystals deposited in the composite have a si milar core shell structure after the preforming step (i.e. a core of  $\alpha$  HQ surrounded by a shell of guest free  $\beta$  HQ).

## 2.2. Apparatus

The experiments were conducted in an HCBGS process at bench scale. This batch process consists of a closed loop across a fixed bed reactor. The main instruments of the setup are: a regulator expander, a Coriolis flowmeter, a gas circulator pump, a reactor, a gas chromato graph, and two coaxial heat exchangers. Except for the reactor and the chromatograph, the equipment is mounted on a support frame specially designed for the process. A P&ID of the experimental setup is shown in Fig. 1.

The reactor (from NovaSwiss) is a column 1.0 m high with an inner

diameter of 25.4 mm, consisting of four removable jacketed sections (each 0.25 m high) made of 316L stainless steel. Five perforated grids are used between each section to prevent particles from flying away. The grids at the inlet and outlet of the reactor have pore sizes of 10 µm, and the intermediate grids have pore sizes of 50 µm. The reactor can run experiments at pressures and temperatures up to 1.5 MPa and 473 K. The temperature is measured by means of nine Pt100 probes (with an accuracy of 0.1 K) placed all along the reactor. The tempera ture is maintained at the desired value with a stability of  $\pm 0.2$  K by continuous forced water circulation in the jacket using a thermostatic bath (Polystat 37, Fischer Scientific). The pressure is measured by three 0 30 MPa Keller model PAA35X sensor transducers (with uncertainties of  $\pm$  0.01 MPa), a numerical manometer (model LeoII) and a 0 20 MPa pressure gauge (from Swagelok). Gas sampling is performed through a needle valve located on the reactor. The gas is analyzed by a gas chromatograph which works with a high pressure sampling system (Agilent, model 7820A). It is equipped with a capillary column of 30 m length and 0.32 mm inner diameter, comprising a stationary phase of polystyrene/di vinylbenzene coating 20 µm thick (model HP PLOT Q) and a thermal conductivity detector. The carrier gas is supplied by a hydrogen generator (model NMH2). The chromatograph is calibrated with different CO<sub>2</sub>/CH<sub>4</sub> gas mixtures allowing for quantitative mea surements with a 0.3% relative uncertainty. The retention times for CH<sub>4</sub> and CO<sub>2</sub> are 0.64  $\pm$  0.05 and 0.80  $\pm$  0.05 min respectively.

The setup is supplied with feed gas by means of a pressure regulator expander (model ER5000 from TESCOM). The expander works with an N<sub>2</sub> gas flow at 0.6 MPa as the pneumatic supply. The feedback pressure signal comes from the pressure sensor transducer at the reactor head. Gas tanks of variable volume are used to feed the whole system. The pressure at the inlet of the expander is measured by a 0 30 MPa Keller model PA33X sensor transducer (with uncertainties of  $\pm$  0.01 MPa). The whole volume placed upstream of the expander is estimated at 994 cm<sup>3</sup>. A vacuum pump is used to produce a primary vacuum of 0.1 kPa which is measured by means of a numerical manometer (model LeoII) and a 0 0.1 MPa pressure gauge (from Swagelok). The gas cir culator pump (model TFK M1 from Gardner Denver) is used to circulate the gas under pressure in the closed loop to homogenize both its tem perature and composition before each chromatographic analysis. The circulation flowrate is adjusted through a voltage regulator. A Coriolis flowmeter (model CMF025H from SERV Instrumentation) checks both the gas flowrate and the amount of gas added to the whole system. Two home made heat exchangers homogenize the gas temperature in the closed loop. The exchangers use water as heat transfer fluid; they consist of double walled pipes with shell outer diameter of 0.02 m and measure 0.5 and 1.5 m in length, respectively. The safety of the HCBGS process is ensured by a relief valve, an electro pneumatic valve, and a safety enclosure.

#### 2.3. Procedure

At the beginning of the experiment, about 200 g of solid reactive medium (native HO crystals or HO based composites) is loaded into the reactor, and the thermostatic bath temperature is set to 293 K. When the temperature is stable, the whole system is put under vacuum to remove the air initially present. All valves are open except for the valves connected to the vents or the atmosphere (V 1, V 10, V 13 and NV 4 in Fig. 1). As the reactor is under vacuum, valve V 12 is closed and the vacuum pump turned off. In addition, the by pass valves (V 7, V 9 and HTV 2), the expander exhaust valve V 4, and the valve V 6 are closed. Thanks to the regulator expander, the system is first pressurized by helium at 0.5, 1.5 and 2.5 MPa to determine the accurate volume of the system (i.e. total volume of the piping and the reactor after the loading of the reactive medium). Then, the thermostatic bath tem perature is set to the desired value and the system is evacuated by opening the valves connected to the vent, and put again under vacuum as described previously. At this point, the system is pressurized by a CO<sub>2</sub>/CH<sub>4</sub> gas mixture at 0.03 MPa per second passing through the Coriolis flowmeter to check the amount of gas added to the system. When the system is at the chosen pressure, valve V 3 is closed, the expander is turned off and V 6 is opened. In addition to the flowmeter measurement, the amount of gas added to the system is determined from Eq. (1):

$$n_{gas}|_{t^{0}} = \frac{PV}{zR(T + \Delta T)}$$
(1)

where z is the compressibility factor (calculated using the Soave Red lich Kwong equation of state and the van der Waals mixing rules with a binary interaction parameter set to 0.1) [35], P is the pressure of the system, T the temperature, V the volume of the gas phase, and  $\Delta T$  the temperature increase due to exothermic effect related to gas loading. The relative deviation between the two determinations of the initial gas amount (i.e. from the flowmeter and the equation of state) is about 5%.

For the isothermal condition of capture, the pressure of the system is tracked as a function of time. The pressure decrease (point A to B in

Fig. 2) is due to gas capture by the reactive medium (either native HQ or HQ based composite). The time for the capture step is set to 96 h (4 days). During the capture step, the gas phase is sampled and ana lyzed by chromatography as shown in Fig. 3. Before each gas compo sition analysis, the gas is homogenized by the circulator pump.

These measurements allow determine, by mass balance, the amount of gas enclosed in the clathrate and its composition. Indeed, the de crease in the molar quantity of the gas phase corresponds to the molar quantity of gas enclosed in the medium. This amount of gas is given by Eq. (2):

$$n_{\text{medium}}^{i}|_{t} = n_{\text{gas}}^{i}|_{t^{0}} - n_{\text{gas}}^{i}|_{t} = \frac{y^{i}PV}{zR(T + \Delta T)} \left|_{t^{0}} - \frac{y^{i}PV}{zRT}\right|_{t}$$
(2)

where superscript i corresponds to constituent  $CO_2$  or  $CH_4$ ,  $y^i$  is the molar composition of constituent i in the gas mixture. The  $CO_2$  mole fraction in the stored gas ( $x^{CO2}$ ), the main determined criterion, can be obtained by the ratio of the amount of stored  $CO_2$  to the total amount of stored gas.

To take into consideration a process point of view, three other cri teria are evaluated: (i) the separation factor (S.F. $^{CO2/CH4}$ ), (ii) the CO<sub>2</sub> recovery fraction (R.F. $^{CO2}$ ), and (iii) the transient storage capacity (q). The latter criterion is the ratio of the amount of gas stored in the clathrate (given by Eq. (2)) to the initial mass of reactive medium.

S.F.  $^{CO2/CH4}$ , which describes the CO<sub>2</sub> capture selectivity, considers the composition of the stored gas in equilibrium with the gas phase. The higher S.F.  $^{CO2/CH4}$ , the better the CO<sub>2</sub> separation. This parameter is calculated by Eq. (3):

S.F.<sup>CO2/CH4</sup>|<sub>t</sub> = 
$$\frac{x^{CO2/1} - x^{CO2}}{y^{CO2} - y^{CO2}}\Big|_{t}$$
 (3)

where  $x^{CO2}$  and  $y^{CO2}$  are the CO<sub>2</sub> molar compositions in the CO<sub>2</sub>/CH<sub>4</sub> gas mixture at time t in the gas stored in the medium and in the gas phase, respectively. R.F.<sup>CO2</sup>, which gives the proportion of CO<sub>2</sub> re covered from the gas mixture initially loaded in the reactor, is obtained by Eq. (4):

$$R.F.^{CO2}|_{t} = 1 - \frac{y^{CO2} \cdot n_{gas}|_{t}}{y^{CO2} \cdot n_{gas}|_{t^{0}}}$$
(4)

For the release step, the system is first depressurized down to 0.1 MPa through the vent (Fig. 2 from point B to C): valve V 11 is closed and valves V 10 and NV 3 are opened (see Fig. 1). The system is then put under vacuum: valves NV 3 and V 10 are closed, valves V 12 and NV 5 are opened, and the vacuum pump is turned on. Once the primary vacuum is reached, valves V 12 and NV 5 are closed, the vacuum pump is turned off, and the thermostatic bath temperature is set to 343 K. The operating conditions of the release step are set to insure the dissociation of the HQ clathrates in agreement with literature data [32]. At this point, the pressure of the system slowly increases due to the gas re leased by the reactive medium until it reaches a constant value (Fig. 2, point D). The composition of the released gas is determined by gas chromatography. The system is again depressurized down to vacuum for further release, followed by composition measurements (if the re active medium has captured enough gas to do this). The release step is completed when the pressure remains invariant over time at a value close to 1 kPa (i.e. primary vacuum) (Fig. 2, point E). During the release step, the composition of the analyzed gas corresponds to the composi tion of the stored gas. The relative deviation of the mass balance cal culated between the capture and release step is less than 1%.

In the experiments, the uncertainties were estimated considering the accuracy of the sensors, the measures of dispersion, and the experimental reproducibility calculated by performing two runs at 293 K/3.0 MPa using the equimolar CO<sub>2</sub>/CH<sub>4</sub> gas mixture.

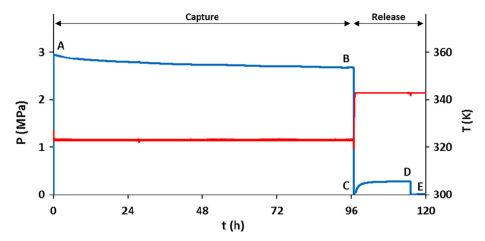


Fig. 2. Example for the capture/release experiment by the HCBGS process loaded with native HQ: system pressure (blue line), and temperature (red line) as a function of time. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

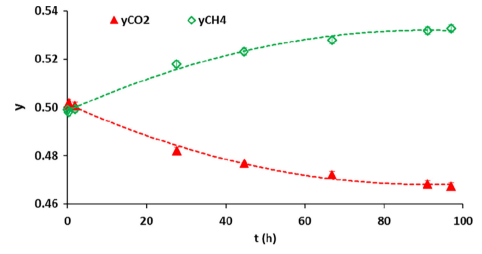


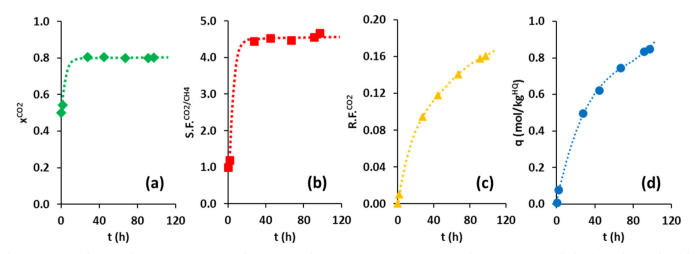
Fig. 3. Example for the capture/release experiment by the HCBGS process loaded with native HQ:  $CO_2$  mole fraction (red line) and  $CH_4$  mole fraction (green line) as a function of time. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

## 3. Results and discussion

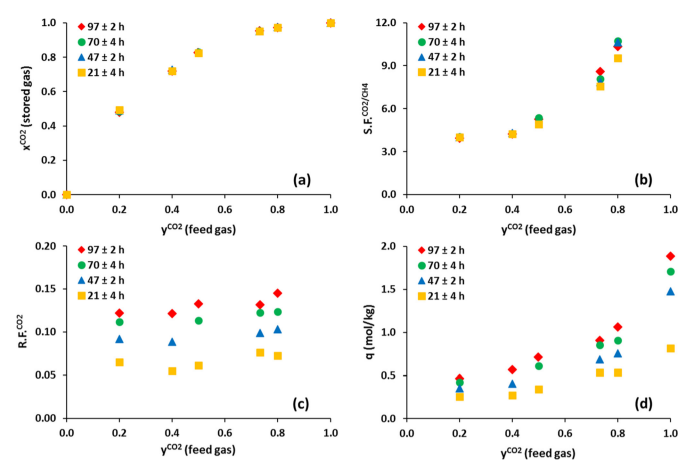
## 3.1. Evaluation of native HQ

First, for each experiment, x<sup>CO2</sup>, S.F.<sup>CO2/CH4</sup>, R.F.<sup>CO2</sup> and q are

tracked as a function of time to assess the reaction progress. Fig. 4 shows an example of this tracking from the equimolar  $CO_2/CH_4$  gas mixture at 323 K and 3.0 MPa. Note that this trend was observed for all pressure and temperature conditions investigated with the native HQ. Using this reactive medium, it appears that the  $x^{CO2}$  stabilizes at a value



**Fig. 4.** Experimental curves of native HQ capturing  $CO_2$  from an equimolar  $CO_2/CH_4$  gas mixture at 323 K and 3.0 MPa: (a)  $CO_2$  mole fraction in the stored gas, (b)  $CO_2$  capture selectivity, (c)  $CO_2$  recovery fraction, (d) transient gas storage capacity as a function of time.



**Fig. 5.** Experimental curves of native HQ capturing CO<sub>2</sub> from different CO<sub>2</sub>/CH<sub>4</sub> gas mixtures at 323 K and 3.0 MPa: (a) CO<sub>2</sub> mole fraction in the clathrate, (b) CO<sub>2</sub> capture selectivity, (c) CO<sub>2</sub> recovery fraction, and (d) transient gas storage capacity. (i)  $21 \pm 4h$ , (a)  $47 \pm 2h$ , (i)  $70 \pm 4h$ , and (i)  $97 \pm 2h$ .

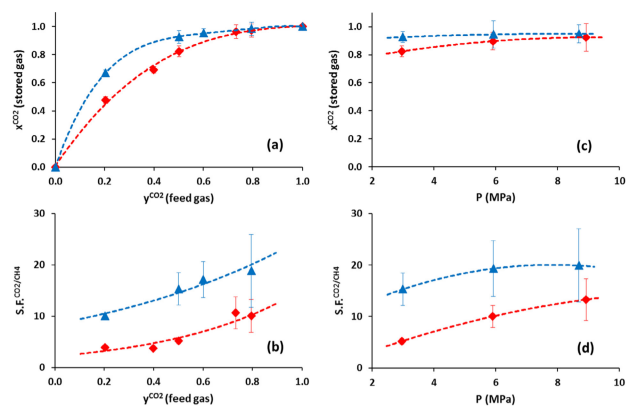
of 0.81 in less than 24 h (Fig. 4 a). Therefore, after a transient period (shorter than 24 h), the preferential gas capture phenomenon by HQ is not a time dependent process. In the same way, after the first day of reaction, the S.F.<sup>CO2/CH4</sup> is almost time independent (Fig. 4 b), as the composition of the gas phase does not change much (i.e. decrease of the CO<sub>2</sub> mole fraction of the gas mixture less than 6%). As the enclathration reaction is not conducted until the full conversion and equilibrium of HQ, the variations of the R.F.<sup>CO2</sup> (Fig. 4 c) and the q (Fig. 4 d) over time are directly linked to the duration of the experiment and are re presentative to the enclathration kinetics.

The effects of the feed gas composition and of the pressure and temperature conditions are then assessed. All the numerical values are given in Table S1, in Supplementary material. The experiments are performed with the different  $CO_2/CH_4$  gas mixtures at 293 and 323 K for pressures of 3.0, 6.0 and 9.0 MPa. The release step is not discussed in this part, as the  $x^{CO2}$  measured during this step is found to be com parable, to within 0.1%, to the one determined in the capture step.

To begin with, we verified the influence of the feed gas composition on the gas capture. Fig. 5 presents  $x^{CO2}$ , S.F. $^{CO2/CH4}$ , R.F. $^{CO2}$  and q for the different CO<sub>2</sub>/CH<sub>4</sub> gas mixtures. In agreement with the previous discussion concerning the time dependence of these criteria, a global trend is observed for all investigated CO<sub>2</sub>/CH<sub>4</sub> gas mixtures: whatever the initial feed gas composition, the preferential capture of CO<sub>2</sub> by native HQ is not time dependent after the first day of reaction (Fig. 5 a). We observed that both the  $x^{CO2}$  and the S.F. $^{CO2/CH4}$  increase with the CO<sub>2</sub> mole fraction in the feed gas (Fig. 5 a, b). Indeed, within the range of compositions investigated at 3.0 MPa and 323 K, the selectivity varies from 3.7 to 10.6. Furthermore, these data indicate that the HQ always preferentially capture the CO<sub>2</sub> contained in the CO<sub>2</sub>/CH<sub>4</sub> mix tures (i.e. the selectivity is higher than 1) in the entire range of the initial gas composition. Regarding R.F.<sup>CO2</sup> and q (Fig. 5 c, d), these criteria increase with the CO<sub>2</sub> mole fraction in the feed gas. Effectively, after the 4 day reaction with feed gas containing 0.2 0.8 of CO<sub>2</sub> mole fraction, R.F.<sup>CO2</sup> increases from 0.12 to 0.14 and q increases from 0.41 to 0.98 mol/kg. Moreover, comparing the latter values against that obtained for a 4 day experiment in the same pressure and temperature conditions with pure CO<sub>2</sub> at about 1.76 mol/kg (Fig. 5 d), we can infer that the presence of CH<sub>4</sub> in the mixture significantly impedes the cap ture kinetics. These observations confirm that the gas capture kinetics is enhanced for high CO<sub>2</sub> mole fractions, as already reported in literature [32].

Hereafter, we consider the criteria obtained for different pressure and temperature conditions after the 4 day reaction (Fig. 6). The ex periments were performed at 293 K over the entire range of the feed gas composition, reaching higher S.F.<sup>CO2/CH4</sup> values than at 323 K (Fig. 6 a, b). The selectivity varies from 10.1 to 18.8 at 293 K, instead of from 3.7 to 10.6 at 323 K. In the reaction between native HQ and the equi molar CO<sub>2</sub>/CH<sub>4</sub> gas mixture (Fig. 6 c, d),  $x^{CO2}$  and S.F.  $CO2/CH4}$  increase at high pressure. When the pressure rises from 3.0 to 9.0 MPa, the se lectivity increases from 5.2 to 13.3 at 323 K, and from 15.3 to 19.9 at 293 K. It is worth noting that the impact of temperature is more im portant than that of pressure: from 3.0 MPa and 323 K, the decrease of 30 K (from 323 to 293 K) increases the S.F. CO2/CH4 by a factor of 3, whereas the pressure increase of 6.0 MPa (from 3.0 to 9.0 MPa) in creases S.F. <sup>CO2/CH4</sup> by a factor of 2. Therefore, the preferential capture of CO<sub>2</sub> is enhanced at low temperature, high pressure and for mixtures having high CO<sub>2</sub> content.

The q obtained after the 4 day capture experiments are presented in Fig. 7. Here, we do not consider the R.F.<sup>CO2</sup> for the comparative pur pose, as the whole volume is not the same between the experiments at

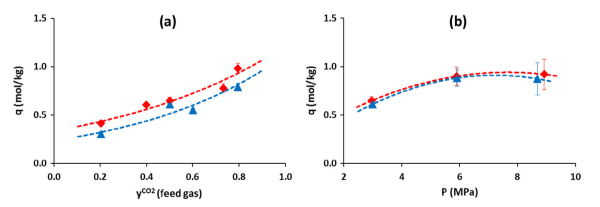


**Fig. 6.** 4-day capture experiments with native HQ at (blue line) 293 K and (red line) 323 K: (left row) for different  $CO_2/CH_4$  gas mixture at 3.0 MPa, and (right row) for the equimolar  $CO_2/CH_4$  gas mixture at 3.0, 6.0 and 9.0 MPa. (a, c)  $CO_2$  mole fraction in the clathrate, (b, d)  $CO_2$  capture selectivity. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

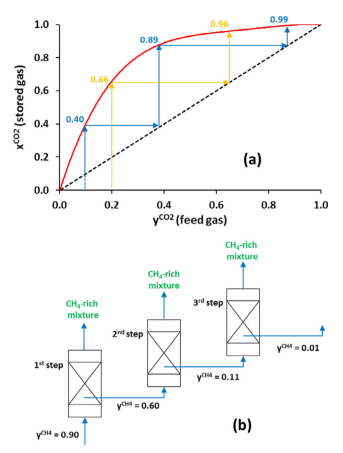
293 and 323 K (i.e.  $677 \pm 14 \text{ cm}^3$  and 723  $\pm 15 \text{ cm}^3$  for the experiments at 293 and 323 K respectively) due to the replacement of the reactive medium in the reactor. Note that the medium used for the experiments comes from the same batch of HQ. The change in gas volume is related to the amount of medium loaded into the reactor which can differ slightly from one experiment to another. Fig. 7 a clearly shows that the kinetics is enhanced for mixture having high CO<sub>2</sub> content whatever the temperature (i.e. 293 or 323 K). As an example, after the 4 day reaction at 293 K and 3.0 MPa, it is found 0.30  $\pm$  0.01 mol/kg for a feed gas mixture having a CO<sub>2</sub> mole fraction of 0.2, whereas the transient capacity is 0.79  $\pm$  0.04 mol/kg for a feed gas mixture having a CO<sub>2</sub> mole fraction of 0.8. Moreover, from Fig. 7 b, the transient gas storage capacity increases with the pressure. Indeed, by increasing the pressure by 6.0 MPa (experiments at 3.0 and 9.0 MPa), the q measured increases by about 50% after the 4 day experiments. The kinetics is thus

improved at high pressure, which is in agreement with other studies [36]. However, it is worth noting that any increase above 6.0 MPa (e.g. from 6.0 to 9.0 MPa) has no significant impact on q. It is possible that the increase in transient capacity related to the enhancement of gas capture kinetics with pressure is affected by a reduction in clathrate occupancy at high pressure as observed in literature [37]. Now, re garding the effect of temperature, we perceive that the high tempera ture slightly enhances the kinetics in agreement with literature data [30]. Interestingly, it can be observed that the variation in temperature has an opposite effect on some of the parameters evaluated in this work: whereas high temperature enhances the enclathration kinetics (and thus the quantity of stored gas during the 4 day experiments), it is detrimental for selectivity (which is enhanced at low temperature).

For the operating point at 293 K and 3.0 MPa, the HCBGS process was conceptualized by means of the McCabe Thiele graphical method



**Fig. 7.** Transient gas storage capacity after 4 days of reaction between native HQ at 293 K (blue line) and at 323 K (red line) and either (a) different  $CO_2/CH_4$  gas mixtures at 3.0 MPa, or (b) the equimolar  $CO_2/CH_4$  gas mixture at 3.0, 6.0 and 9.0 MPa. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



**Fig. 8.** Conceptual HCBGS process. (a) McCabe-Thiele graphical method: (red line)  $CO_2$  mole fraction in the clathrates as a function of  $CO_2$  mole fraction in the feed gas at 293 K and 3.0 MPa, and (dotted line) x = y line. (b) Schematic diagram of a three-stage HCBGS process at 293 K and 3.0 MPa. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

(Fig. 8). The experimental results are used in a polynomial fit to represent the  $CO_2$  mole fraction in the clathrates as a function of  $CO_2$  mole fraction in the feed gas. As our study is directly related to gas sweetening, we illustrate the HCBGS process through two realistic ex amples using feed gases with a  $CO_2$  mole fraction of 0.2 and 0.1. It thus appears that the  $CO_2$  mole fractions of the  $CO_2/CH_4$  gas mixtures can be concentrated to 0.96 and 0.89 through a two stage HCBGS process from feed gases with a  $CO_2$  mole fraction of 0.2 and 0.1, respectively. Fur thermore, a third stage could enrich the recovered mixture to almost pure  $CO_2$  (i.e.  $CO_2$  mole fraction of the  $CO_2/CH_4$  gas mixture of 0.99).

## 3.2. Evaluation of the HQ based composite

Now considering the HQ based composite, we first investigated the adsorption properties of the native silica support, as the composite is known for combining both adsorption on silica and enclathration by HQ [27]. Table 1 gives the quantity of gas adsorbed and the selectivity obtained for the adsorption of the equimolar  $CO_2/CH_4$  gas mixture at 283, 293, 323 and 343 K for 3.0 and 6.0 MPa on native silica particles.

In all the investigated conditions, the quantity of gas adsorbed on the silica particles is very low (at most 0.16  $\pm$  0.04 mol/kg at 283 K and 6.0 MPa): this represents between 1.3 and 5.2% of the maximum gas storage capacity of the HQ clathrates (i.e. 3.03 mol/kg<sup>HQ</sup>). This trend is effectively in line with the high pore size (i.e. macro pores of 100 nm) and the low specific area (i.e. specific area of 57 m²/g) of these particles. However, as for the CO<sub>2</sub> capture selectivity by native silica, it is clear that CO<sub>2</sub> is preferentially adsorbed on this medium in all the

#### Table 1

Quantity of gas adsorbed and selectivity of SiliaSphere<sup>®</sup> silica particles in contact with an equimolar  $CO_2/CH_4$  gas mixture.

P (MPa)	T (K)	q <sub>ads</sub> (mol/kg)	S.F. <sup>CO2/CH4</sup>
$\begin{array}{c} 3.00 \pm 0.02 \\ 3.00 \pm 0.02 \\ 3.00 \pm 0.02 \\ 2.99 \pm 0.02 \\ 5.99 \pm 0.02 \\ 6.00 \pm 0.02 \\ 6.00 \pm 0.02 \\ 6.03 \pm 0.02 \end{array}$	$283.2 \pm 0.2  293.2 \pm 0.2  323.2 \pm 0.3  343.2 \pm 0.3  283.2 \pm 0.2  293.2 \pm 0.2  323.2 \pm 0.3  343.2 \pm 0.3  343.2 + 0.3  $	$\begin{array}{c} 0.11 \ \pm \ 0.03 \\ 0.08 \ \pm \ 0.02 \\ 0.06 \ \pm \ 0.02 \\ 0.04 \ \pm \ 0.03 \\ 0.16 \ \pm \ 0.04 \\ 0.15 \ \pm \ 0.04 \\ 0.08 \ \pm \ 0.02 \\ 0.06 \ \pm \ 0.02 \end{array}$	$32 \pm 17 \\ 28 \pm 14 \\ 6 \pm 2 \\ 2 \pm 1 \\ 39 \pm 20 \\ 9 \pm 5 \\ 6 \pm 3 \\ 3 + 2 $

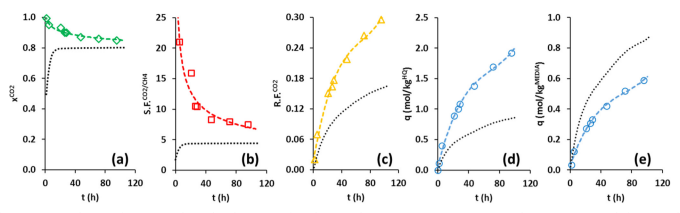
tested pressure and temperature conditions (i.e. S.F.<sup>CO2/CH4</sup> > 1). Moreover, we observed that the selectivity is higher at low temperature, and considering the uncertainties, it is probable that selectivity remains unchanged for the investigated pressures. The general trend known for adsorption capacity on silica is observed: enhancement at high pressure and low temperature [38 39].

From measurements involving the HQ based composite (Fig. 9), we observe that x<sup>CO2</sup> decreases versus time, as does S.F.<sup>CO2/CH4</sup> (Fig. 9 a, b). These observations indicate that initially there is a fast, strong capture of  $CO_2$  molecules followed by a second capture that is less  $CO_2$ selective and slower than the first. The fall in the selectivity over time cannot be explained by the two successive phenomena of adsorption on silica and enclathration by HQ. Indeed, the selectivity is about 22 at the start of the experiment for the HQ based composite at 323 K and 3.0 MPa, whereas the native silica achieves a selectivity of 6 in the same conditions. However, as discussed previously in the section on "Mate rials", we believe that after the dissociation step, most of the HQ crystals have a core shell structure composed of a core of preformed  $\alpha$ HO surrounded by a shell of guest free  $\beta$  HQ. We have already de monstrated in a previous study that the HQ guest free structure formed with  $CO_2$  is much more selective to  $CO_2$  than preformed  $\alpha$  HQ [32]. Accordingly, the highest selectivity values measured with the compo site just after contacting the gas mixture to HO could be explained by the presence of the guest free  $\beta$  HQ structure. As the contact area of the impregnated HQ (i.e. crystallites sized 100 nm at most) is much higher than that of native HQ, it would be expected to contain a greater pro portion of guest free  $\beta$  HQ and thus have a greater effect on selectivity

in the HQ based composite. We believe that the same gas capture mechanisms by enclathration through HQ [32] occur both with the native crystals and with the composite. Nevertheless, as the proportion of guest free structures is supposed to be much higher in the composite than in the native HQ crystals (proportional to the surface to volume ratio of the HQ in the two reactive media), the effects on selectivity are much more visible when the composite is used.

Additionally, Fig. 9 b shows that after 4 days of reaction, the se lectivity appears slightly higher for the HQ based composite in com parison to pure HQ. This is logical as the composite allows coupling of the adsorption on silica, filling of the guest free  $\beta$  HQ, and capture by  $\alpha$  HQ, in which the first two phenomena are more CO<sub>2</sub> selective than typical enclathration from native HQ. Interestingly, such result suggests that different HQ clathrates may coexist: the final state is composed of CO<sub>2</sub> rich clathrates (obtained from the guest free) and HQ clathrates containing a lower quantity of CO<sub>2</sub> (obtained from the  $\alpha$  HQ).

Now, as regarding R.F.<sup>CO2</sup> (Fig. 9 c), it is obvious that more  $CO_2$  can be recovered from the gas phase by the HQ based composite compared against the native HQ. Indeed, the composite doubles the R.F.<sup>CO2</sup>. Moreover, the quantity of gas captured by the impregnated HQ from the composite materials (i.e. transient gas storage capacity of the clathrate) is found to be higher than that of the native HQ after the 4 day reaction (Fig. 9 d). This confirms that the composite material greatly improves the enclathration kinetics [26]. Nevertheless, it is worth noting (e.g. for scaling up the process) that due to the high



**Fig. 9.** Experimental curves of  $CO_2$  capture by HQ-based composite, from the equimolar  $CO_2/CH_4$  gas mixture at 323 K and 3.0 MPa: (a)  $CO_2$  mole fraction in the stored gas, (b)  $CO_2$  capture selectivity, (c)  $CO_2$  recovery fraction, (d) transient gas storage capacity of the clathrate, (e) transient gas storage capacity of the media, as a function of time. Black dotted lines represent the values obtained for native HQ in the same pressure and temperature conditions.

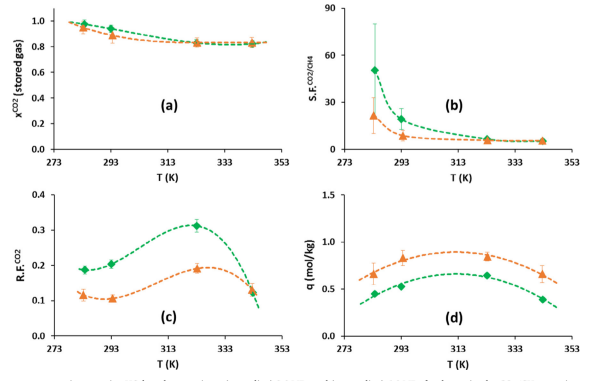
proportion of silica in the composite material, the transient global storage capacity of this medium (i.e. molar quantity of  $CO_2$  stored by reference to the mass of the composite) is lower than for native HQ (Fig. 9 e) (e.g. after the 4 day reaction at 323 K and 3.0 MPa, the quantity of gas captured by native HQ is about 20% higher than for the composite material).

The HQ silica composites were then tested in the 4 day capture experiments with the equimolar  $CO_2/CH_4$  gas mixture in the tempera ture range of 283 343 K, and at pressures of 3.0 and 6.0 MPa. x<sup>CO2</sup>, S.F.<sup>CO2/CH4</sup>, R.F.<sup>CO2</sup> and q are shown in Fig. 10. All the numerical values are given, for convenience, in Table S2, in Supplementary material. From the evolution of x<sup>CO2</sup> and S.F.<sup>CO2/CH4</sup> as a function of temperature (Fig. 10 a, b), the same trend is observed as that previously discussed for native HQ. However, as for the effect of pressure (experiments at 3.0 and 6.0 MPa) and according to the uncertainties, it is worth noting that these parameters have a negligible impact on the selectivity.

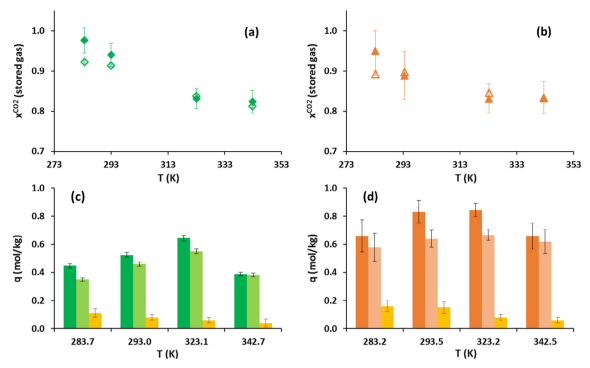
Concerning the evolution of R.F.<sup>CO2</sup> and q as a function of

temperature (Fig. 10 c, d), we observed that the maximum value could be reached for temperatures between 293 and 323 K. As the different capture phenomena occurring with the composite materials (i.e. adsorption on silica, filling of the guest free  $\beta$  HQ, and capture by the  $\alpha$  HQ) vary in the same way with temperature, the presence of such extrema cannot be linked to them. Therefore, in the temperature range investigated, it appears that both low temperature (< 293 K) and high temperature (> 323 K) have an adverse effect on gas capture by the HQ based composite. However, enclathration being the main phe nomenon, and knowing that the storage capacity of HQ clathrates (at equilibrium) is enhanced at low temperature [30], the extremum found on the curves shown in Fig. 10 d for moderate temperature is then lo gical.

The influence of pressure on the recovery fraction (see Fig. 10 c) is in line with the  $CO_2$  capture selectivity, which is somewhat lowered with this parameter. However, the transient storage capacity increases



**Fig. 10.** 4-day capture experiments using HQ-based composite at (green line) 3.0 MPa and (orange line) 6.0 MPa for the equimolar  $CO_2/CH_4$  gas mixture at 283, 293, 323 and 343 K: (a)  $CO_2$  mole fraction in the clathrate, (b)  $CO_2$  capture selectivity, (b)  $CO_2$  recovery fraction, (d) transient gas storage capacity. The volume of the system is 696 ± 25 cm<sup>3</sup>. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



**Fig. 11.** 4-day capture experiments using HQ-based composite at (a, c) 3.0 MPa and (b, d) 6.0 MPa for the equimolar  $CO_2/CH_4$  gas mixture at 283, 293, 323 and 343 K. (a, b)  $CO_2$  mole fraction of the stored gas: (full symbols) capture step and (open symbols) release step. (c, d) transient gas storage capacity measured: (dark color) capture step, (light color) release step, and (yellow) quantity of gas adsorbed on the native silica particles. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

by about 30% when the pressure increases from 3.0 to 6.0 MPa (Fig. 10 d).

and native silica.

4. Conclusions

When the composite is used, we noted a difference in the mass balance between the gas capture and release steps. As a reminder, in the experimental protocol the release step begins by putting the reactor under vacuum. The gas adsorbed on the support of the composite (i.e. the silica) is evacuated in this step, and cannot be directly measured when the vacuum pump is turned on. Therefore, we assumed that the totality of the gas released after the reactor was put under vacuum is the gas contained in only the HQ clathrates. This assumption is supported by the fact that the clathrate dissociation is rather slow even in vacuum conditions [27,32]. Accordingly, two other parameters that are re lated only to the HQ clathrates and specific to the release step can be defined and estimated: the CO<sub>2</sub> mole fraction in the clathrates ( $x_c^{CO2}$ ) and the quantity of gas captured in the clathrate. The results are pre sented in Fig. 11 and the values are given in Table S2.

For the experiments performed at low temperature, the  $x_{\rm c}^{\rm CO2}$  mea sured in the release step (Fig. 11 a, b) is found to be generally lower than the  $x^{\text{CO2}}$  determined in the capture step (e.g. at 283 K, the  $\text{CO}_2$ mole fraction are 0.976  $\pm$  0.031 and 0.923  $\pm$  0.003 for the capture and the release step, respectively). However, at high temperature the two values match to within about 1% (e.g. at 323 K, the CO2 mole fraction values are 0.834  $\pm$  0.080 and 0.833  $\pm$  0.002 for the capture and the release step, respectively). Fig. 11 c, d also presents the transient gas storage capacity determined in both the capture and re lease steps. As observed, the quantity determined in the capture step is always higher than that in the release step, confirming that adsorption on the support always occurs. Interestingly, the difference between these two determinations is of the same order of magnitude as the quantity of gas adsorbed on the native silica. This is logical, as even though part of the silica surface is covered by HQ crystals in the com posite materials, most of the surface is available for adsorption [27]. Therefore, it appears that the HQ based composite materials allow for efficient coupling of adsorption and enclathration: the resulting synergy implies that this medium can be more interesting than both native HQ

## This study presents an assessment of the separation of the $CO_2/CH_4$ gas mixture through a hydroquinone clathrate based gas separation (HCBGS) process. $CO_2$ is selectively captured when either native HQ or HQ based composite materials contacts a $CO_2/CH_4$ gas mixture with a $CO_2$ mole fraction from 0.2 to 1, in temperatures ranging of 283 343 K and pressures from 3.0 to 9.0 MPa.

Using native HQ, it is demonstrated that the preferential capture of  $CO_2$  molecules is not time dependent after about 24 h, and is enhanced at low temperature, high pressure and for mixtures with a high  $CO_2$  content. After 4 day capture experiments and for all the conditions investigated, the quantity of gas captured in this period (i.e. the tran sient storage capacity) varies in the same way as selectivity, whereas the gas capture kinetics is enhanced at high temperature, high pressure and for gas mixtures with a high  $CO_2$  content. With the apparatus used and the time interval set for the capture step (4 days in this work), a two stage HCBGS process would be sufficient treating a  $CO_2/CH_4$  gas mixture with a  $CO_2$  mole fraction of at least 0.1, as it is possible to achieve concentrated gas with a  $CO_2$  mole fraction of 0.89.

Concerning the HQ silica composite material, the coupling of ad sorption and enclathration phenomena is confirmed. Furthermore, we demonstrated a high  $CO_2$  selective phenomenon occurring at the be ginning of the reaction between the composite and the  $CO_2/CH_4$  gas mixtures, attributed to the filling of a substantial amount of guest free  $\beta$  HQ present in the reactive medium. The  $CO_2$  capture selectivity of the composite can thus be preferred over the capture selectivity of either the native HQ or native silica. Moreover, experiments involving the composite confirm that the enclathration kinetic is greatly improved by using this reactive medium. Nevertheless, and despite the kinetics gain, the high proportion of silica contained in HQ based composite materials implies that the global capacity (i.e. molar quantity of  $CO_2$  stored by reference to the mass of the composite) is lower compared to native HQ. Finally, from a process point of view, the HCBGS process requires certain compromises with the operating parameters, and the three in dicators that define the separation efficiency (i.e. the capture selectivity toward CO<sub>2</sub>, the kinetics of gas capture, and the transient gas storage capacity of the material used) cannot all be maximized together using a single set of parameters: (i) low temperature conditions insure both high selectivity and high capacity, whereas high temperatures favor the capture kinetics; (ii) the use of an HQ based composite insures high selectivity to  $CO_2$  and enhances the gas capture kinetics, while the global capacity is reduced compared to pure HQ.

Therefore, taking into consideration all the information available on thermodynamics (clathrate formation can be achieved at low pressure and high temperature levels), kinetics (technical solutions are proposed to improve the enclathration rate), and selectivity (preferential capture of  $CO_2$ ), the HCBGS process could be a valuable alternative to the conventional gas separation processes and may seriously surpass hy drate based technologies for this gas mixture. Additional work is now in progress in our laboratory to evaluate the performance of the HCBGS process for other gas mixtures.

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## **Author Contributions**

The manuscript was written by contributions from all authors. All authors have given approval to the final version of the manuscript.

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#### Notes

The authors declare no competing financial interest.

## Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.fuel.2018.03.170.

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# Hydroquinone Clathrate based Gas Separation (HCBGS): Application to the CO<sub>2</sub>/CH<sub>4</sub> Gas Mixture.

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# SUPPLEMENTARY MATERIAL

# 1. EVALUATION OF THE NATIVE HQ

Table S1. 4-days capture experiments with  $CO_2/CH_4$  gas mixtures and native HQ.

V (cm <sup>3</sup> )	P (MPa)	T (K)	y <sup>CO2</sup> (feed gas)	x <sup>CO2</sup> (stored gas)	S.F. <sup>CO2/CH4</sup>	<b>R.F.</b> <sup>CO2</sup>	q (mol/kg)	v <sup>Gas</sup> (STP)/v <sup>HQ</sup>
723 ± 15	$2.97\pm0.02$	$323.2 \pm 0.3$	$0.204 \pm 0.001$	$0.477 \pm 0.023$	3.9 ± 0.2	$0.12 \pm 0.01$	0.41 ± 0.03	12.0 ± 0.9
$723 \pm 15$	$3.01\pm0.02$	$323.3 \pm 0.3$	$0.399\pm0.001$	$0.692 \pm 0.024$	$3.7 \pm 0.2$	$0.12 \pm 0.01$	$0.61\pm0.02$	$17.6\pm0.7$
723 ± 15	$2.96\pm0.02$	$323.2 \pm 0.3$	$0.501 \pm 0.002$	$0.824 \pm 0.040$	$5.2 \pm 0.6$	0.13 ± 0.01	$0.65 \pm 0.04$	18.9 ± 1.1
$723 \pm 15$	$2.95 \pm 0.02$	$323.2 \pm 0.3$	$0.733 \pm 0.002$	$0.963 \pm 0.048$	$10.1 \pm 3.1$	0.13 ± 0.01	$0.78 \pm 0.04$	$22.7 \pm 1.3$
723 ± 15	$2.97 \pm 0.02$	$323.3 \pm 0.3$	$0.795 \pm 0.002$	$0.972 \pm 0.046$	10.6 ± 3.2	$0.14 \pm 0.01$	$0.98 \pm 0.05$	28.7 ± 1.5
$677 \pm 14$	$2.98\pm0.02$	$293.2 \pm 0.3$	$0.204 \pm 0.001$	$0.671 \pm 0.020$	$10.1 \pm 0.4$	$0.23 \pm 0.01$	$0.30 \pm 0.01$	8.9 ± 0.3
$677 \pm 14$	$2.99\pm0.02$	$293.2 \pm 0.3$	$0.501 \pm 0.002$	$0.926 \pm 0.045$	$15.3 \pm 3.1$	$0.20 \pm 0.02$	$0.61 \pm 0.04$	$17.8 \pm 1.2$
$677 \pm 14$	$2.99\pm0.02$	$293.2 \pm 0.3$	$0.601 \pm 0.002$	$0.955 \pm 0.029$	$17.1 \pm 3.5$	$0.19 \pm 0.01$	$0.55 \pm 0.02$	$16.1 \pm 0.5$
$677 \pm 14$	$2.97 \pm 0.02$	$293.5 \pm 0.3$	$0.795 \pm 0.002$	$0.984 \pm 0.046$	18.8 ± 7.1	$0.16 \pm 0.01$	0.79 ± 0.04	23.0 ± 1.2
$682 \pm 14$	$5.91 \pm 0.02$	$323.1 \pm 0.5$	$0.501 \pm 0.002$	$0.897 \pm 0.063$	$10.0 \pm 2.1$	$0.15 \pm 0.02$	$0.90\pm0.10$	$26.2 \pm 2.8$
$682 \pm 14$	$5.92 \pm 0.02$	$293.6 \pm 0.3$	$0.501 \pm 0.002$	$0.945 \pm 0.062$	19.3 ± 5.4	$0.12 \pm 0.01$	$0.88 \pm 0.09$	$25.7 \pm 2.6$
$715 \pm 14$	$8.92 \pm 0.03$	$323.0 \pm 0.6$	$0.501 \pm 0.002$	$0.924 \pm 0.098$	$13.3 \pm 4.0$	$0.10 \pm 0.01$	$0.92 \pm 0.16$	$26.8 \pm 4.6$
715 ± 14	8.69 ± 0.03	$293.5 \pm 0.3$	$0.501 \pm 0.002$	$0.949\pm0.107$	19.9 ± 7.1	$0.08 \pm 0.01$	$0.87 \pm 0.17$	25.5 ± 4.9

# 2. EVALUATION OF THE HQ-BASED COMPOSITE MATERIALS

**Table S2.** 4-days capture experiments with the equimolar  $CO_2/CH_4$  gas mixtures and the HQ-basedcomposite. The volume of the system is  $696 \pm 25 \text{ cm}^3$ .

P (MPa)	T (K)	x <sup>CO2</sup> Capt. step (stored gas)	S.F. <sup>CO2/CH4</sup>	<b>R.F.</b> <sup>CO2</sup>	q <sup>Capt. Step</sup> (mol/kg)	x <sup>CO2</sup> Rel. step (stored gas)	q <sup>ReL step</sup> (mol/kg)
$2.99\pm0.02$	$283.7\pm0.4$	$0.976 \pm 0.031$	50.5 ± 29.5	$0.19\pm0.01$	$0.45\pm0.02$	$0.923 \pm 0.003$	$0.35\pm0.01$
$2.98\pm0.02$	$293.0\pm0.3$	$0.940 \pm 0.029$	$19.3 \pm 6.7$	$0.20\pm0.01$	$0.52\pm0.02$	$0.913 \pm 0.003$	$0.46\pm0.02$
$3.00\pm0.02$	$323.1 \pm 0.4$	$0.831 \pm 0.025$	6.6 ± 1.0	$0.31\pm0.02$	$0.64\pm0.02$	$0.838 \pm 0.003$	$0.55\pm0.02$
$3.02\pm0.03$	$342.7\pm0.8$	$0.824 \pm 0.028$	$5.2 \pm 0.9$	$0.12 \pm 0.01$	$0.39\pm0.02$	$0.812 \pm 0.002$	$0.38\pm0.01$
6.03 ± 0.03	$283.2 \pm 0.4$	$0.951 \pm 0.050$	$21.5 \pm 11.4$	$0.12 \pm 0.02$	$0.66 \pm 0.11$	$0.898 \pm 0.003$	$0.58\pm0.10$
$5.99\pm0.02$	$293.5\pm0.3$	$0.889 \pm 0.059$	8.8 ± 3.4	$0.11\pm0.01$	$0.83\pm0.08$	$0.897 \pm 0.003$	$0.64\pm0.06$
$6.05\pm0.02$	$323.2 \pm 0.4$	$0.831 \pm 0.036$	$5.8 \pm 1.2$	$0.19\pm0.01$	$0.84\pm0.05$	$0.846 \pm 0.003$	$0.67\pm0.03$
$6.00\pm0.02$	$342.5 \pm 1.1$	$0.834 \pm 0.040$	5.6 ± 1.3	$0.13\pm0.02$	$0.66 \pm 0.09$	$0.833 \pm 0.002$	$0.62 \pm 0.09$