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Hydrate growth at the interface between water and pure or mixed CO₂/CH₄ gases: Influence of pressure, temperature, gas composition and water-soluble surfactants

Delphine Daniel-David, Fabrice Guerton, Christophe Dicharry, Jean-Philippe Torré, Daniel Broseta

Laboratoire des Fluides Complexes et leurs Réservoirs Univ Pau and Pays Adour, 64013 Pau Cedex, France

- Hydrate growth is observed from experiments with water drops and CH₄/CO₂ gases.
- Hydrates most often form low permeable crusts at water/gas interfaces.
- Anionic surfactants such as SDS or AOT promote CH₄ hydrates but not CO₂ hydrates.
- Insight is given into the capillary driven hydrate growth observed with SDS and AOT.

Key words: CO₂/CH₄ hydrates
Hydrate growth
Hydrate promoters
Surfactants

1. Introduction

Clathrate hydrates (hereafter abbreviated as hydrates) consist of hydrogen bonded water (host) molecules forming a crystalline lattice stabilized by hydrate former (guest) molecules present in some of its cavities. These solid (ice like) compounds are stable for temperatures and pressures usually higher than the water/ice transition temperature (0°C) and the atmospheric pressure. Since most hydrate formers are sparingly soluble in water, hydrates usually appear and grow at/from the interface between the (liquid or gas) guest phase and the water rich phase. The formation, growth and morphology of hydrates at guest/water interfaces under quiescent conditions are the focus of increasing interest, motivated both by fundamental reasons and practical applications (Chatti et al., 2005; Sloan and Koh, 2008).

On the fundamental side, the respective roles of heat and mass transfers (Mochizuki and Mori, 2006; Saito et al., 2010; Sun et al., 2010) and of surfactant molecules added in small amounts to the water or guest phases (Zhong and Rogers, 2000; Karanjkar et al., 2012; Mitarai et al., in press) are not yet fully understood. When the hydrate former is sparingly soluble in water, hydrate formation occurs at the water/guest interface, along which a
polycrystalline thin crust grows rapidly (Taylor et al., 2007). This thin crust contains gas filled pores that anneal over time, which increases resistance to mass transfer (Davies et al., 2010) and results in a very slow crust thickening and growth normal to the interface: over reasonable timescales and under quiescent conditions, much of the water trapped beneath this hydrate crust remains unconverted (Sloan and Koh, 2008). In some (rare) instances, however, the growing porous structure, rather than forming a low permeable thin crust at the guest/water interface, allows much of the water (if guest molecules are in excess) to be rapidly and totally converted into hydrate. This still ill understood hydrate formation mechanism (often referred to as ‘capillary driven’) is encountered for instance with methane and low molecular weight alkanes (but not with CO₂) when anionic surfactant additives such as sodium alkyl sulfates (e.g., sodium dodecyl sulfate or SDS) are added to the water in small amounts (typically, a few hundreds of ppm) (Zhong and Rogers 2000; Sun et al., 2003a, b; Lin et al., 2004; Gayet et al., 2005; Okutani et al., 2008).

On the side of practical applications, the ability to speed up hydrate formation and growth is key to the success of emerging hydrate based technologies such as natural gas or hydrogen storage and transportation by means of hydrates, refrigeration processes using hydrates as a phase change material, CO₂ separation, water purification and desalination, etc. Clearly, a capillary driven mechanism (promoted by proper surfactant additives) of hydrate formation is preferred for these applications, unless a large amount of guest/water interfaces is generated by, e.g., mechanical agitation, spraying/bubbling one phase into the other (Gnanendran and Amin, 2004; Brinchi et al., 2014), or circulating the guest phase through mesoporous particles saturated with water (Dicharry et al., 2013). In other applications such as oil and gas transport through pipelines, hydrate formation and growth must be impeded to ensure flow, which is achieved by using low dosage (a few hundreds to thousands of ppm) water soluble molecules called kinetic hydrate inhibitors (KHi). The performance of these KHi might be related to their ability to slow down the lateral growth of the hydrate crust at the water/guest interface where they usually adsorb (Peng et al., 2009; Duchateau et al., 2012; Wu et al., 2013); another important factor seems to be the localization of hydrate growth at the interface, which impedes or delays the growth of a 3D porous structure by the capillary driven flow mechanism mentioned above.

This paper presents an experimental investigation of hydrate growth at the interface between water and mixtures of CO₂ and CH₄ as a function of temperature, pressure, gas composition and surfactant additives present in the water phase. The following aspects are examined: hydrate crust morphologies and lateral growth rates, and capillary driven growth in the presence of some surfactant additives.

Studies on hydrate growth at the interface between the aqueous and guest phases have recently been reviewed by Sun et al. (2010), who did not however consider water/guest systems in which the guest phase is a mixture of gases and/or an aqueous phase containing a small amount of surfactant additives. We therefore first briefly review the work done on these systems, after reviewing for completeness that on pure water/CO₂ and CH₄ systems.

Uchida et al. (1999) were the first to report observations of CO₂ hydrate growing on the surface of water droplet suspended in CO₂ and noticed that the propagation rate is primarily dependent on, and an increasing function of, subcooling ΔT = Tₑq - Tₑx where Tₑx is the temperature of the experiment and Tₑq the dissociation temperature (i.e., the temperature of hydrate/water/gas equilibrium) at the pressure of the experiment. A similar dependence was observed by Freer et al. (2001) for CH₄ hydrate growing at a water/CH₄ (planar) interface, albeit at a rate lower by about one order of magnitude than CO₂ hydrate at similar subcooling. These and other authors (Servio and Englezos, 2003; Ohmura et al., 2004) examined hydrate crust morphologies: a more faceted aspect (individual crystals of millimeter sizes) is observed at low subcooling ΔT, while a smooth appearance is noted at intermediate and high ΔT. Extensive observations of hydrate crust texture and lateral growth at water/gas interfaces have been conducted over the past decade, most of them focused on water/CH₄ systems, using various configurations: a rising (gas) bubble in water (Peng et al., 2007; Sun et al., 2007; Li et al., 2013, 2014), a water drop (in gas) either pendant (Zhong et al., 2011) or sitting on a substrate (Tanaka et al., 2009), or a planar water/gas interface (Kitamura and Mori, 2013). While there is some scatter in the published lateral growth rates, all observations show that the hydrate crust texture gets smoother with increasing subcooling and increasing time. The variation of lateral growth rates with subcooling is accounted for by models in which heat transfer processes at the edge of the advancing hydrate front is the controlling factor (Mochizuki and Mori, 2006; Sun et al., 2010), but mass transfer limitations appear to play an important role as well (Saito et al., 2010).

A few studies have been conducted with gas mixtures as the guest phase or with some water soluble additives present in the water phase. The measurements by Peng et al. (2007) of hydrate growth at the surface of bubbles (in pure water) of various CH₄+C₃H₈ mixtures, as well as one CH₄+C₅H₁₂ mixture, indicate lateral growth rates smaller than those of the corresponding pure water/pure gas systems. A similar effect has been observed with CH₃-C₅H₁₂ mixtures by Li et al. (2014), who argued that the coexistence in the hydrate crust of two crystalline structures (I and II) might be responsible for the slowing down of the frontal advance. However, the latter effect was not observed in the experiments conducted by Saito et al. (2011) on sessile water drops with two CH₄ rich (90+ mol. %) mixtures of CH₄+C₅H₁₂ and C₂H₆. The studies conducted with surfactant or polymeric additives present in the water phase are of two sorts. One is concerned with polymeric additives inhibiting hydrate formation (KHs), which delay nucleation and/or slow down hydrate film lateral growth at water/guest interfaces (Peng et al., 2009; Duchateau et al., 2012; Wu et al., 2013). The other is concerned with additives (such as SDS) promoting hydrate formation. The understanding of these promoting processes, loosely referred to as ‘capillary driven’, is one of the most challenging fundamental issues in gas hydrate research, whereas on a more practical side simple and rapid methods are needed for assessing the hydrate promoting potential of a given additive. As stated by Lo et al. (2012), finding the efficient surfactant(s) among the hundreds of existing surfactants is currently like ‘finding a needle in a hay stack’: quick assessment methods such as those proposed by these authors, or the drop based method proposed in this paper, are urgently needed.

The outline is as follows: The next section describes the gases and surfactant additives used, as well as the apparatus and experimental methodology for forming and visualizing hydrate growth on/in a water drop in a gas atmosphere at controlled pressure and temperature. Then, in Section 3, the results obtained with pure water are presented and discussed: the effects of gas composition, temperature and pressure on hydrate crust texture and growth rate are analysed in detail. Section 4 is devoted to the results obtained under the same conditions as in Section 3, with the difference that a small amount of surfactant is present in the water phase: some insights are given into capillary driven hydrate formation.

2. Experiments

2.1. Materials

CO₂ and CH₄ were purchased from Linde (99.995%) and CO₂/CH₄ mixtures from Air Liquide (see Table 1 for compositions).
Table 1
Compositions of the CO₂/CH₄ gas mixtures used in the experiments (as given by Air Liquide).

<table>
<thead>
<tr>
<th>No. of mixed gas sample</th>
<th>CO₂, mol%</th>
<th>CH₄, mol%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>q.s</td>
<td>24.98 ± 0.50</td>
</tr>
<tr>
<td>2</td>
<td>q.s</td>
<td>49.00 ± 0.99</td>
</tr>
<tr>
<td>3</td>
<td>25.08 ± 0.50</td>
<td>q.s</td>
</tr>
</tbody>
</table>

Those compounds and mixtures were used in the experiments under monophasic (gas) conditions, i.e., at low enough pressures in order to avoid the condensation of the CO₂ rich mixtures at the temperatures of interest (from 0 to 5 °C). Water is deionized and has a resistivity of 18 MΩ cm.

2.2. Surfactant additives

2.2.1. Non ionic additives

The non ionic surfactants, chosen for their reported ability to lower the interfacial tension and/or to promote emulsions between water and dense (supercritical) CO₂, belong to four different classes. Some of them have been already tested in the context of gas hydrate research (Sun et al., 2003a, b; Zhang et al., 2008; Zhong et al., 2008; Delahaye et al., 2011; Dickrey et al., 2013). The first class of surfactants consists of polyoxyethylene sorbitan fatty acid esters, with commercial names Tween® plus a number specifying the fatty acid: 20 (monolaurate), 40 (monopalmitate), 60 (monostearate) and 80 (monoooleate). The second class consists of polyethylene oxide/propylene oxide triblock or random copolymers with commercial names Pluronics®: they are called by a letter (for the physical state: L for liquid, P for paste) and a number for the molecular weight (first one or two digits) and the proportion of ethylene oxide monomers (last digit) and in between the letter R if the copolymer is random. The third class of surfactants consists of polyethylene oxide 2,6,8 trimethyl 4 nonyl ethers known as Tergitol® TMNn, where n(=6 and 10 in the present study) is the number of ethylene oxide monomers. Lastly, an ethoxylated acetylenic diol with trade name Surfynol 465 has been tested. The characteristics (supplier, molecular weight, hydrophilic lipophilic balance or HLB) and molecular formulae of the surfactants tested are listed respectively in Table S1 and Figure S1 of Supplementary material.

2.2.2. Anionic surfactants

The anionic surfactants tested (only with pure CH₄ and pure CO₂) are SDS (sodium dodecyl sulfate) dissolved in water at concentra tions of 100 and 500 ppmw (i.e., 0.01 and 0.05 wt%), and a common Gemini (double tailed) surfactant, dioctyl sodium sulfocinate (also known as AOT), at a concentration of 100 ppmw. SDS is chosen because this is the prototypical CH₄ hydrate promoter. Gemini surfactants need a lower dosage than SDS for a given surface tension decrease and activity at water/hydrate interfaces (Salako et al., 2013). Some of these surfactants have been observed to give comparable formation rates and higher methane storage capacities than SDS at very low dosage (below 150 ppm) (Kwon et al., 2011).

2.2.3. Interfacial behavior

The interfacial tensions between the aqueous and gas phases were measured in the course of the drop experiments (see below) at the experimental temperatures (from 0 to 5 °C) and pressures (from 27 to 40 bar); the contact angles of a drop of the aqueous solutions on PTFE, a strongly hydrophobic substrate, were measured as well (see Section 2.3 for the measurement method). The above surfactants indeed lowered water/guest interfacial tensions and rendered the substrate less hydrophobic, but in limited manner for most solutions (see Table S1 in Supplementary material). All tensions remained above 20 mN/m and contact angles (measured in the aqueous phase) above 50–60°, except for the 1000 ppmw Tergitol® TMN6 and TMN10 solutions that exhibited (see Table S2) lower tensions and angles in presence of CO₂ (no measurement was attempted with the other gases).

2.3. Experimental apparatus and procedure

The experiments of hydrate growth at the water/gas interface of a water drop are conducted under controlled temperature and pressure. Hydrate formation is triggered at the top of a sessile water drop (in a gas atmosphere) by contact with another hydrate hanging at the tip of a capillary, then the subsequent hydrate growth is monitored by means of an image acquisition and treatment system. The procedure is similar to that used by Peng et al. (2007), Sun et al. (2007) and Li et al. (2014), with the difference that these researchers nucleated the hydrate at the top a gas bubble (immersed in the aqueous phase) by contact with another gas bubble covered with hydrate.

The experimental apparatus, schematically presented in Fig. 1, consists of

(i) A high pressure (maximum operating pressure: 20 MPa) viewing cell (volume 17 cm³) with two parallel see through sapphire windows of 18 mm diameter, designed in collaboration with Top Industrie (France). The cell contains a horizontal plate and a vertically moveable capillary tube connected to a Teledyne ISCO Syringe Pump (100 DM) allowing a drop of the aqueous phase to be formed at the tip of the capillary and eventually be deposited on the horizontal plate. The cell and plate temperatures are controlled independently by means of two refrigerating/heating circulation systems (Fischer Scientific Polystat 37). The temperature in the cell is measured by means of a Pt100 sensor with an accuracy of 0.1 °C.

(ii) An optical and imaging system, composed of a diffuse light source, a CCD camera (Toshiba 8 bits; maximum acquisition speed: 25 fps; sensor size: 6.5 mm × 4.85 mm, 752 × 582 pixels, 680 × 480 of which are exploited in the image treatment software) and a PC. The image treatment software software (Windrop™, Teclis, Longessaigne, France) provides the contour of the sessile drop for each image and computes the drop area and volume, as well as the contact angle and interfacial tension (the contour is analyzed in terms of the Young Laplace equation, provided volume densities are given for details see Broseta et al., 2012).

The experimental procedure comprises the following steps:

![Fig. 1. Schematic diagram of the experimental apparatus. (1) high-pressure cell, (2) pendant drop, (3) capillary tube, (4) sessile drop, (5) temperature-controlled plate, (6) CCD camera, (7) computer, (8) gas storage vessel, (9) syringe pump.](image-url)
(1) Gas is flushed into the cell and evacuated three times in order to evacuate air.

(2) The cell is filled with gas at the operating pressure at ambient temperature.

(3) A drop of the aqueous phase is deposited by means of the vertically moving capillary tube on the horizontal plate previously (unless otherwise specified) covered with a PTFE film. In all measurements presented here, the area of this sessile water drop, denoted \( a \), lies in the range of \( 40 \pm 8 \text{ mm}^2 \).

(4) The cell temperature is then lowered to \( 3 \text{ °C} \) and gas is injected into the cell to compensate for the pressure decrease. Spontaneous hydrate nucleation may occur at the surface of the sessile water drop: in this case, the experiment is restarted and cell temperature is brought to, e.g., \( 2 \text{ °C} \).

(5) Gas hydrate (to be used for triggering hydrate formation by contact with the top of the sessile drop, see step 6) is formed at the tip of the capillary as follows: a pendant water drop formed at the tip of the capillary is first pumped into the upper part of the capillary, which is cooled down to about \( 30 \text{ °C} \) by contact with the immersion probe of a Julabo FT401 immersion cooler, then the pump pushes back to the tip of the capillary some crystallized water that reacts with the ambient gas to form hydrates (Fig. 2(a)). The sessile water drop (see step 3) is still fully liquid.

(6) Temperature is raised to the desired value, and the system left to equilibrate for a while. The hydrate crystal hanging at the tip of the capillary (see step 5) is then brought in contact with the top of liquid sessile drop by moving the capillary downwards (Fig. 2(b)), which triggers hydrate formation and subsequent growth. In most situations, hydrate growth consists in the propagation of a hydrate crust at the surface of the sessile water drop starting from the contact point to the base of the sessile water drop. This process is recorded with the CCD camera until there is no detectable change or movement. One of the recorded movies (corresponding to the snapshots of Fig. 2) is available in the supplementary information (movie M1). Two types of information are extracted from these movies:

Supplementary material related to this article can be found online at http://dx.doi.org/10.1016/j.ces.2015.04.015.

The morphology (or texture) of the gas hydrate crust formed at the interface between water and gas and its evolution with time, the growth rate of the hydrate crust, defined as an average surface (or two dimensional) rate \( r_a \) equal to the sessile drop area \( a \) divided by the time \( t_r \) required for the drop to appear as ‘frozen’ or immobile (see Appendix A) or as an average linear rate \( r_b \) equal to the drop contour divided by \( 2 \times t_r \) (the hydrate crust travels from the top to the base of the drop); following Sun et al. (2007), surface rates \( r_a \) are reported here, and linear rates are evaluated only with pure CO\(_2\) and pure CH\(_4\) for the purpose of comparison with published results.

The plate is heated for several minutes in order to dissociate the hydrate on the sessile water drop, while the cell remains at the operating temperature; some crystallized gas hydrate remains hanging at the tip of the capillary previously raised to the upper part of the cell (see step 6).

Then, a new measurement is started by bringing the crystallized gas hydrate in contact with the sessile water drop (step 6, etc.).

A minimum of four measurements is carried out for each temperature, pressure, gas and aqueous phase composition, from which the average growth rate is calculated, together with the related standard deviation. The results are reported in Table 2 (pure water) and S2 in Supplementary material.

The independent temperature control of the plate is very practical for carrying out numerous consecutive runs and for assessing the reproducibility of growth rate measurements. No gas consumption is detected during experiments due to the very small volume of hydrate formed by comparison with the cell volume. The features of the CCD camera allow taking pictures with a resolution of 640 X 480 pixels and a frame rate of 30 frames per second, which is sufficient for the present experiments.

Fig. 2. Snapshots of CO\(_2\)-hydrate growth on a sessile water drop (at 26.6 bar, 5 °C) showing the approach of the capillary with a hydrate phase at its extremity (a), contact of this hydrate with the top of the water drop (b), propagation of the hydrate crust along the surface of the drop (c) and final aspect of the drop (d). Duration of crust propagation is about 15 s. Drop diameter (white arrow) is equal to 4.5 mm. (See movie M1, Supplementary material.)
camera limit the range of pressures and temperatures investigated: for high driving forces (i.e., high pressure and low temperatures), growth is too rapid to be detected by the CCD camera.

3. Results and discussion: experiments conducted with pure water

The drop experiments carried out with pure water, CO2, CH4, and various CO2/CH4 gas mixtures (with compositions given in Table 1) are presented first. They have been conducted at temperatures $T_{\text{exp}}$ in the range of 0–5°C and pressures $P_{\text{exp}}$, not exceeding 45 bar in the case of pure CH4 and 27 bar in the case of pure CO2 (and intermediate values for the gas mixtures), in such a way that the non aequous phase remains gaseous. Unless otherwise specified, each line in Table 2 corresponds to a set of (at least four) experiments conducted with the same gas composition, pressure and temperature conditions (columns 1–3). The subcooling $\Delta T=T_{\text{eq}}-T_{\text{exp}}$, where $T_{\text{eq}}$ is the dissociation temperature at $P_{\text{eq}}$ ($T_{\text{eq}}$ is determined by using the software CSMGem, Sloan and Koh, 2008) and the dimensionless driving force (see Appendix B) are reported in columns 4 and 5. (For the conditions investigated here, the dimensionless driving force is proportional to the subcooling $\Delta T$, in agreement with previous results on similar systems, see Arjmandi et al., 2005). Finally, the average of the measured growth rates $r_\text{a}$ and the associated standard deviations are reported in the rightmost column of Table 2, together with the linear rates $r_\text{l}$ (in the case of pure CO2 and pure CH4 only).

In the case of pure CH4, the hydrate crusts formed on the water drop surface are barely visible (they are smoother and thinner, see next section): the results corresponding to only three different temperature and pressure conditions are reported in Table 2.

The results displayed in Table 2 clearly indicate that the growth of the hydrate crust is more rapid at higher subcooling $\Delta T$ and higher CO2 content in the guest gas. Morphological aspects are examined prior to presenting and discussing growth rates in a quantitative fashion.

3.1. Hydrate morphologies

The morphology (or texture) of the hydrate crust covering the water drop is observed to vary with gas composition and driving force (or subcooling). This morphology experiences some changes after the drop has been fully covered with hydrate, as commented below.

3.2. Effect of gas composition and driving force (or subcooling)

Fig. 3 depicts the final aspect of a water drop covered with hydrate crusts formed at similar subcooling (3–4°C) but different compositions ranging from 100% CO2 to 0% CH4. Hydrate layers of pure methane (and methane rich) hydrates are smooth and their presence at the drop surface barely alters the shape of the initial water drop, whereas CO2 hydrate crusts show well defined crystals (in the micron range) and thus looks smooth under moderate magnification, whereas at low driving force (below $\Delta T=1$ K in the case of CH4 hydrates) the crystallites get large (a few tens to hundreds of μm) faceted or polygonal shapes.

In many experiments, changes are still occurring after the water drop has been fully covered with hydrate. In some instances, a second hydrate layer seems to follow the formation of the first hydrate crust, resulting in a smoother surface, or in the smoothing of an initially textured crust: see Fig. 5 that depicts how the hydrate crust formed from pure water and a 25 mol% CO2/75 mol% CH4 gas mixture has its vertical stripes smoothed out within about one second (this smoothing proceeds from the top to the bottom of the drop in such a way that it looks like a second hydrate crust is propagating). A possible mechanism is as follows: the thinner regions of the initially irregular crust are more permeable to gas and water and therefore get thicker more rapidly. This phenomenon has been observed by Uchida et al. (1999) at the interface between water and CO2, and by Zhong et al. (2011) at the interface between water and CH4.

3.3. Lateral growth rates

The measured lateral growth rates $r_\text{a}$ given in Table 2 are displayed as a function of subcooling $\Delta T$ in Fig. 6 for the pure CH4 and CO2 and the three CH4/CO2 mixtures investigated. For a given
guest gas, these rates strongly increase with subcooling as a power of $\Delta T$ with an exponent in the range of 2 (see Fig. 6) in agreement with many observations (Peng et al., 2007; Sun et al., 2007) and with a theoretical approach based on heat transfer considerations (that predicts an exponent equal to 2.5, see Mochizuki and Mori, 2006; Peng et al., 2007). A good fit of the data is also provided by the exponential correlation proposed by Sun et al. (2007),

$$r_g = A \exp\left(\frac{B \Delta g}{RT}\right).$$

where $\Delta g/RT$ is the dimensionless driving force (see Appendix B). For a given subcooling $\Delta T$, growth rates strongly increase with the fraction of CO2 in the guest gas. For example, at $\Delta T \approx 5.5 \, ^\circ C$, growth rates increase by more than one order of magnitude, from $2.7 \, \text{mm}^2 \, \text{s}^{-1}$ (pure methane) to nearly $40 \, \text{mm}^2 \, \text{s}^{-1}$ (pure CO2). This increase is strongly non linear: it is moderate (less than a 2 fold increase) when the CO2 molar content increases from 0% (pure CH4) to 25% and to 50%, and more pronounced when the CO2 content increases from 50% to 75% and to 100% (see Fig. 6).

Our three data points on CH4 hydrate growth rates can straightforwardly be compared to those of Sun et al. (2007), who observed for dimensionless driving forces from 0.37 to 0.48 growth rates increasing from 1.0 to 1.7 mm$^2$ s$^{-1}$, slightly lower than our values from 1.4 to 2.7 mm$^2$ s$^{-1}$. This difference might be ascribed to the different configuration and procedures: a quicker removal of the heat generated by hydrate formation is ensured in our experiments, where the configuration is that of a sessile water drop in a gas environment, whereas that of Sun et al. (2007) is that of a gas bubble immersed in water; in addition, the sessile water drop is placed onto a cooled plate in our procedure, whereas only the cell as a whole is temperature controlled in the experiments by Sun et al. (2007).

The measured linear rates $r_l$ (rightmost column in Table 2) are in line with those observed with CO2 on water drops by Uchida et al. (1999), but somewhat higher than those observed (again with CO2) by Peng et al. (2007) on gas bubbles. With CH4 our results are in line with those observed by Peng et al. (2007) and by Li et al. (2014) on gas bubbles, and by Freer et al. (2001) on planar water/gas interfaces, but about twice higher than those observed by Kitamura and Mori (2013), also on planar water/gas interfaces.

To explain the observed large difference in lateral growth rates for hydrates formed from CO2 (and CO2 rich) gases and from CH4 (and CH4 rich) gases, other effects than heat transfer processes have to be taken into account, as thermal properties of these two systems are very similar. Gas solubility in water evidently plays an important role. Ohmura et al. (2004) were the first to build a model in which gas solubility (or, more precisely, the gas solubility difference between the hydrate free and hydrate containing water) controls the kinetics of hydrate growth, and later established a correlation between lateral growth rate and gas solubility (Saito et al., 2010). The monotonic trend with gas composition that we observe differs from the trend observed by Peng et al. (2007) with mixed CH4/C2H6 gases and by Li et al. (2014) with mixed CH4/C2H4 gases, who noted lower rates for hydrates formed from gas mixtures in comparison to the rates of the corresponding pure gas hydrates. As mentioned in the Introduction, Li et al. (2014) attributed this trend to the formation of hydrates of a different structure (II) for some mixed CH4/C2H6 gases.
4. Results and discussion: experiments conducted with watersoluble additives

We present in this section the results of the experiments conducted with systems and conditions similar to those described in the previous section: the only difference is the composition of the aqueous phase, to which is added a small concentration (in the range of 100 to 1000 ppm) of non ionic or anionic surfactant molecules.

4.1. Non ionic surfactants

At least one surfactant of each of the four classes of non ionic surfactants (Tweens®, Pluronics®, Tergitols®, and Surlyn®s, see Section 2) has been tested with the pure gases CO₂ and CH₄ and the three CO₂/CH₄ mixtures listed in Table 1, the largest number of surfactants being tested with pure CO₂. Surfactant concentration is equal to 100 ppmw in most experiments, and to 500 or 1000 ppmw in a few cases. For a given gas composition, the temperatures and pressures conditions are similar to those used with pure water (see preceding section).

All experiments conducted with a non ionic surfactant as water soluble additive do exhibit the standard growth pattern that is observed in the absence of additives, that is, rapid lateral growth of a hydrate crust at the interface between the aqueous and the gas phases, and growth rates are not very different from those observed in the absence of additive (see preceding section): rate values are reported in Table S2 (Supplementary material). However, morphologies may differ from one additive to the other: an example is given in Fig. 7, where spherule like structures are apparent at the surface of a water drop containing 1000 ppmw of Pluronics (P104).

4.2. Anionic surfactants: SDS and AOT

With pure CO₂ those two surfactants (with concentration in water equal to 100 ppmw) have little effect on hydrate growth on the drop surface, which again obeys the ‘standard’ growth pattern and exhibits rate values similar to those observed in the absence of additives (see Table S2).

With pure CH₄, however, the observed hydrate growth pattern depends on both the nature of the anionic surfactant and its concentration. For the lowest SDS concentration investigated (100 ppmw) at 0 °C and 40 bar, the ‘standard’ hydrate growth pattern is observed, with a lateral growth rate somewhat higher than with pure water; 1.4 vs. 2.4 mm² s⁻¹ (Table S2). For the highest SDS concentration investigated (500 ppmw) at the same T (0 °C) and P (40 bar), ‘capillary driven’ growth is observed: following contact at its top with the hanging hydrate (formed from a 500 ppmw SDS solution), the sessile drop is rapidly ‘swallowed’ by the hanging hydrate, which grows in size as it incorporates more water (Fig. 8 and movie M2 in Supplementary material). In this growth process all the water contained in the sessile drop is transformed into a solid hydrate phase (the gas phase being in large excess): the rightmost picture in Fig. 8,
where the hanging hydrate has been broken in two pieces by contact with the lower substrate, reveals the internal solid-like structure (see also movie M2 in Supplementary material).

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For the 100 ppmw AOT solution, a growth behavior is observed that is close to that observed with the 500 ppmw SDS solution, even though it proceeds more slowly and ends up in a somewhat different morphology: most of the sessile aqueous drop is ‘swallowed’ into the hanging hydrate (also formed from a 100 ppmw AOT solution), which undergoes a gradual and slow (within 5-6 min) transformation into a grape-like structure (Fig. 9).

A strong initial wetting of the hanging CH$_4$ hydrate by the aqueous phase is apparent in the experiments involving SDS and AOT. This is particularly apparent when the hanging hydrate hits the sessile drop of aqueous solution, which immediately imbibes what looks like a porous solid-like structure (see Movie M2, Supplementary material). The analogous experiments conducted with CO$_2$ do not exhibit such a strong apparent initial wetting (of the hanging CO$_2$ hydrate by the aqueous SDS or AOT solution).

The capillary driven character of gas hydrate formation is usually characterized from macroscopic measurements, e.g., from pressure or temperature variations in large volume cells, but less often from observations of the evolution of the water/gas interface over mesoscopic (sub millimetric) scales. Our results for the 100 ppmw SDS solution agree with those of Sun et al. (2007), who did notice an accelerated lateral growth of the hydrate crust along the surface of CH$_4$ bubbles in such a solution, but our results for the 500 ppmw SDS solution disagree with those by the same authors, who were able to measure lateral growth rates at the surface of CH$_4$ bubbles in a 500 ppmw SDS solution. They also differ from the recent observations by Lee et al. (2014) on bubbles of a 90/10 mol% CH$_4$/C$_3$H$_8$ gas mixture in very low concentrated (from 10 to 50 ppm) SDS solutions; these authors observed halos (of small hydrate crystals?) erupting and rising above the immersed gas bubble, which however preserved its integrity. Our results of the Gemini AOT surfactant (compared to that of SDS) as regards CH$_4$ hydrate formation complement those by Kwon et al. (2011) and by Salako et al., 2013, who point to a superior performance of Gemini surfactants.

One question that arises from the above observations is the following. Why is a ‘capillary driven’ hydrate formation process, i.e., a complete and rapid conversion of low concentrated SDS and AOT aqueous solutions into hydrate, occurring with CH$_4$ and not with CO$_2$? This question raised by many authors (starting with Zhang and Lee, 2009) has not received any clearcut answer yet. Some molecularly based mechanisms have recently been invoked (Alberti et al., 2013), but it is likely that larger scale (mesoscopic) mechanisms are (also) at play. First insights into these mesoscale mechanisms are gained by noting the analogy between hydrate crystals growing at the aqueous surface and the salt efflorescences that grow at the evaporative surface of a porous medium saturated with brine. These efflorescences evolve between two limiting patterns, called crusty and patchy

5. Summary and conclusions

A simple experimental method has been devised that allows a rapid assessment of the potential of a given water soluble surfactant additive in promoting hydrate growth. This method consists in triggering hydrate formation and growth at the top of a sessile water drop by contact with the hydrate phase, and in visualizing the ensuing hydrate growth. When the method is implemented with gases consisting of CH$_4$, CO$_2$ and their mixtures, and water, either pure or containing a low concentration of a non ionic or anionic surfactant, two very different hydrate growth patterns are observed. The most prevalent growth pattern, which occurs in all situations involving pure water or a low concentrated solution of non ionic surfactant, consists in the rapid growth of a low permeable hydrate crust laterally at the water/guest interface, at a rate that increases with subcooling and CO$_2$ content, and depends much less on the nature of the non ionic surfactant, at least in the ranges of low concentrations (100 ppmw, occasionally 500 and 1000 ppmw) and low pressures not exceeding 27 bar for CO$_2$, 45 bar for CH$_4$, and intermediate pressures for CO$_2$/CH$_4$ mixtures investigated. Once these interfaces are fully covered with the hydrate crust, the conversion of water and gas into hydrates is almost stopped. The other, ‘capillary driven’ growth pattern, is observed with methane and anionic surfactants (SDS, AOT) at concentrations in the range of 500 w and 100 ppmw, respectively: a permeable hydrate skeleton forms that continuously ‘pumps’ the aqueous solution and brings it in contact with the hydrate former (methane) until no water is left for conversion.

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Appendix A. Image treatment procedure for determining the water drop coverage time $t_f$

This procedure exploits the following feature: after being covered with hydrate, the water drop appears to be ‘frozen’, i.e., it no longer moves (at least over the time scale of the experiment, i.e., at most a few minutes). This occurs at a time $t_f$ after the drop has been contacted with hydrate at its top. This time, which is well defined because the change to a ‘frozen’ state occurs abruptly, is determined from an automated procedure that first splits the movie into its consecutive images (labeled by integer indexes $i$, each image corresponding to a specified time) and then in analyzing all sets of two consecutive images, starting from the image of first contact between the water drop and the hydrate hanging at the tip of the capillary. Each image contains $320 \times 256$ pixels where each pixel has values from 0 (white) to 255 (black). For each couple of consecutive images (labeled by index $i$ corresponding to the first of the two images), the following quantities are calculated:

(i) The standard deviation of the difference between the two images,
(ii) the correlation coefficient (corr2 in the Matlab™ library) between the two images, equal to 1 for identical images (hence, one minus this coefficient is the quantity considered), and
(iii) the maximum pixel absolute value (on a 0-255 scale) of the difference between the two images.

An example of how those three quantities vary with $i$ (or, equivalently, with elapsed time, as there is an interval of $\sim 0.15$ s between two consecutive images) is shown in Figure A1, which corresponds to the experiment giving the snapshots in Fig. 2 (pure water and CO$_2$ at 5 °C and 26.6 bar).

We observed that, as a rule, the transition between the high values (corresponding to an advancing hydrate film) and the low values due to noise (the fully hydrate covered drop no longer moves) is very sharp and occurs at similar image number (or elapsed time) for the three above quantities, which thus provides an unambiguous determination of $t_f$. The corresponding image number is determined automatically in the procedure as follows.

First, the asymptotic (i.e., at large time or large index $i$) average values and standard deviations of those three quantities are computed over a common range of image indexes (or elapsed time) such that these values appear fairly constant (e.g., from 260 to the maximum image number in the example of Figure A1). The quantity (iii) is prone to larger fluctuations (because it singles out one pixel) than quantities (i) and (ii): this is taken into account in the criterion for deciding when the drop starts being ‘frozen’ through the standard deviation, see below.

Second, the image number is decreased incrementally starting from the largest index $i$, and the above three quantities (i, ii, and iii) are calculated for each corresponding couple of images. When the quantity $i$, ii or iii exceeds for at least three consecutive indexes its asymptotic value plus one standard deviation it is denoted by a vertical line in Figure A1. The indexes of these vertical lines usually coincide or differ very slightly, which permits times $t_f$ to be defined unambiguously, as illustrated in Figure A1.

Appendix B. Calculation of the dimensionless driving force

The driving force at a certain temperature $T$ and pressure $P$ below the equilibrium pressure $P_{eq}$ is defined as $\Delta g$, the molar Gibbs free energy difference between the experimental and equilibrium conditions (the minus sign is to render the expression positive):

$$
\Delta g = \upsilon w P_{eq} + RT \sum x_i \ln \left( \frac{p_{eq}}{p_{exp}} \right)
$$

![Fig. A1. Values of the quantities (i), (ii) and (iii) for each set of two consecutive images as a function of time or image number $i$ (see text). The experiment is that illustrated in Fig. 2. The scale on the vertical axis is arbitrary.](image-url)
\[ \frac{f_{eq}^C}{f_{eq}^H} = \frac{RT}{h_{eq}^i} \sum x_i \ln \left( \frac{f_{eq}^i}{P^{\alpha}_i} \right) + \Delta G(P, P^{\alpha}). \]

where \( f_{eq}^C \) and \( f_{eq}^H \) are the fugacities of component \( i \) (CO\(_2\) and CH\(_4\)) at experimental and equilibrium conditions, and \( x_i \) is the mole fraction of component \( i \) in the gas phase, respectively; they are calculated by the Peng Robinson equation of state. \( h_{eq}^i \) is the molar volume of liquid water, \( v_{eq} \) is the molar volume of water in hydrate and \( \Delta G \) is the dimensionless driving force is the driving force divided by \( RT \), where \( R=8.3145 \text{ J/molK} \) is the perfect gas constant.

### Appendix C. Supplementary data

Supplementary data associated with this article can be found in the online version at [http://dx.doi.org/10.1016/j.ccs.2015.04.015](http://dx.doi.org/10.1016/j.ccs.2015.04.015).

### References


