A study on the influence of nanofluids on gas hydrate formation kinetics and their potential: Application to the CO2 capture process

Samer Said, Varun Govindaraj, Jean-Michel Herri, Yamina Ouabbas, Mohamed Khodja, Mohamed Belloum, Jitendra Sangwai, Ramamurthy Nagarajan

To cite this version:

HAL Id: hal-01346311
https://hal.archives-ouvertes.fr/hal-01346311
Submitted on 21 Jul 2016

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
A study on the influence of nanofluids on gas hydrate formation kinetics and their potential: Application to the CO$_2$ capture process

Said Samer$^{1,*}$, Varun Govindaraj$^2$, Jean-Michel Herri$^3$, Yamina Ouabbes$^3$, Mohamed Khodja$^4$, Mohamed Belloum$^1$, Jitendra S. Sangwai$^5$, Ramamurthy Nagarajan$^2$

$^1$Laboratory of Materials Chemistry and the Living (LCMVAR), University Hadj Lakhdar Batna-1, 05 a.v. Chahid Boukhlouf, 05000 Batna, Algeria

$^2$Department of Chemical Engineering, Indian Institute of Technology Madras, Chennai, Tamil Nadu 600036, India

$^3$Gas Hydrates Dynamic Centre, ENSM Saint-Etienne, 158 cours Fauriel, 42023 Saint-Etienne, France

$^4$Sonatrach-Upstream Activity, Technologies and Development Division, Avenue 01 November 35000 Boumerdes, Algeria

$^5$Gas Hydrate and Flow Assurance Laboratory, Petroleum Engineering Program, Department of Ocean Engineering, Indian Institute of Technology Madras, Chennai, Tamil Nadu 600036, India

Corresponding Author:
Samer said: saidpg2006@gmail.com
Phone: +213 662 38 69 00
ABSTRACT

In this work, the effects of Al$_2$O$_3$, SiO$_2$, Ag and Cu nanoparticles on the kinetics of CO$_2$-CH$_4$ hydrate formation process were experimentally studied by measuring the amount of gas consumed and the rate of gas consumption. A suspension of 0.1wt%, 0.2wt % and 0.3wt% of each nanoparticle was injected into the hydrate formation reactor, while pressure and temperature were maintained at 4.0 MPa and 274.15 K, and the magnetic stirrer speed was set at 350 rpm. The CO$_2$-CH$_4$ hydrate formation process was studied in both pure water and water containing a 0.1wt%, 0.2wt% and 0.3wt% of each nanoparticle suspension. The results showed that these nanoparticles had a positive effect on hydrate formation. These effects varied from one nanoparticle to another. It was observed that nanoparticles of SiO$_2$ had the most positive effect on CO$_2$ gas consumption, particularly at a concentration of 0.3wt%. At this concentration the average amount of gas consumed was about 45% higher than that in pure water during the dissolution and 77% during crystallization. Cu and Al$_2$O$_3$ nanoparticles had an intermediate effect with improvement in gas consumption by 1% to 15% during dissolution; while it had an important impact on gas consumption during hydrate crystallization with an improvement of 30% to 65%. Ag nanoparticles had no significant effect during these two phases.

Keywords: Gas Hydrates; Nanoparticles; Kinetics; CO$_2$.

1. INTRODUCTION

The main techniques used to capture the CO$_2$ at the source are based on absorption processes (chemical or physical), adsorption and membrane methods. These techniques are efficient, but often have high energy costs (nearly USD $20 per ton of CO$_2$) (Deppe et al., 2006). In addition to these conventional methods, there are new separation techniques
currently under investigation which are based on the use of low temperature distillation, ionic liquids, and, more recently, gas hydrate crystallization (Mondal et al., 2012).

CO$_2$ capture process by hydrate formation is proving to be one of the promising options for long term carbon storage. This technology is desirable as CO$_2$ hydrate deposited in the deep ocean where the suitable pressure and temperature is ready without any extra cost. The principle of this process is based on hydrate selectivity towards CO$_2$; the trapping of CO$_2$ molecules is preferred to the trapping of other molecules present in the charge. There are certain barriers that should be overcome before this technique could have the industrial applications. The sluggish process of hydrate formation is one of significant problems in the industrial application; therefore, investigating the kinetics of hydrate formation and searching for effective promoters is very important. Several studies have been carried out on the effects of some surfactants on gas hydrate induction time, growth rate, and storage capacity (Ganji et al., 2007a,b; Zhang et al., 2007; Karimi et al., 2013). The addition of nanoparticles to the aqueous phase is a novel approach for the enhancement of the gas hydrate formation kinetics by increasing heat and mass transfer. There are several methods for creating nanoparticles, including attrition, pyrolysis and hydrothermal synthesis (Sobhani et al., 2014; Ansari et al., 2014; Sobhani et al., 2015). The suspensions of these nanoparticles in base fluids showed enhancement ability in heat and mass transfer so that gas dissolution and heat releasing efficiency increased in comparison with their base fluids (Kim et al., 2007; Özerinç et al., 2010; Torres et al., 2012; Jung et al., 2012; Lee and Kang, 2013).

In 2006, Li et al. (2006) studied the formation and dissociation of HFC134a (CH$_2$FCF$_3$) gas hydrates in the presence of copper nanoparticles. This study was inspired from the positive effect of nanoparticles on transfer phenomena (mass and heat) in fluids. It was conducted over a number of different mass fractions of nanoparticles. The results indicated that the addition of these particles increased both mass and heat transfer during the
process of formation and dissociation of HFC134a hydrates. Park et al. (2010) studied the effect of water based solutions of MWCNTs (Multi-Walled Carbon Nano-tubes) on the formation rate of methane hydrates and on the amount of gas consumed during the formation process. They used pure water as the base fluid, and added to it a dispersion of MWCNTs of 10 to 15 nm in diameter and 10-20 microns in length for different mass fractions (0.001, 0.002, 0.003, 0.004, 0.005 and 0.006 wt%). The results showed an increase in gas consumption and in the rate of formation of methane hydrates (e.g, a nanofluid of 0.004 wt% led to an increase of nearly 300% in the amount of gas consumed during the hydrate formation process, relative to pure water).

Another study by Kim et al. (2011) compared the effect two kinds of MWCNTs (CM-95 and CM-100) had on the methane hydrate formation rate (the length of CM-95 is much smaller than that of CM-100). Pure water was used as the base fluid and different concentrations (0.001, 0.002, 0.003, 0.004, 0.005 and 0.006 wt%) of CM-95 and CM-100 were dispersed in it to form a solution. They found that the maximum gas consumption was observed at a concentration of 0.004 wt% of CM-95 and CM-100. For CM-95, the experiments showed that gas consumption for a weight fraction of 0.004 % exceeded that of pure water by nearly 300%. Also, at the same temperature, the methane hydrate formation pressure was lower for CM-95 in water at 0.004 wt% when compared to CM-100 in water at the same mass fraction. This indicates an unexpected thermodynamic effect. The results also showed that the methane hydrate formation time was significantly reduced when using CM-95 and CM-100 nanofluids at 0.004wt%, more prominently so for the CM-95 nanofluid. They concluded that MWCNTs with short lengths were more effective in reducing the hydrate formation time. A second study that was carried out by Park et al. (2012) was on the characteristics of methane hydrates in two types of carbon nanofluids, namely MWCNT/water nanofluids and OMWCNT/water nanofluids (oxidized MWCNT). They
chemically oxidized the surface of the MWCNT and then dispersed it in distilled water. They found that the amount of gas consumed during the formation of methane hydrates in nanofluids of OMWCNT/water were 4.5 times higher than that in nanofluids of MWCNT/water.

In 2013, Ganji et al. (2013) investigated the effect of polymers and polymer nanoparticles suspensions on stability and storage capacity of methane gas hydrate. They found that some tested additives can increase the stability of the formed hydrate effectively. The hydrate formed in the presence of these additives had higher gas content compared to the pure water hydrate. Another study by Mohammadi et al. (2014) focused on the kinetic formation of CO₂ hydrates in the presence of silver nanoparticles and a surfactant (sodium dodecyl sulfate, SDS). These systems were tested in a stirred reactor, with an aqueous solution of SDS at concentrations of 300 and 500 ppm, nanofluids of Ag at concentrations of 0.000045 M and 0.00009 M, and in a mixture of SDS (500 ppm) and nanoparticles of Ag (0.000045 M). The obtained results showed that individually, the SDS and Ag nanoparticles did not have a significant effect on the induction time and hydrate storage capacity, however, the mixture of SDS and Ag nanoparticles led to an increase in the CO₂ storage capacity.

The effects of the graphite nanoparticles on the CO₂ hydrate formation process was studied by Zhou et al (2014). The results showed that the nanographite particles had a positive effect on the CO₂ hydrate formation. In comparison to pure water, the induction time of CO₂ hydrate formed in the presence of the graphite nanoparticles decreased by 80.8%, while the maximum CO₂ consumption increased by 12.8%. In addition, the hydrate reaction was 98.8% complete within 400 min in the presence of nanographite particles. Mohammadi et al. (2016) studied the effect of synthesized zinc oxide (ZnO) nanoparticles on the kinetic and thermodynamic equilibrium conditions of CO₂ hydrate formation. The amount of the gas
consumption was measured and compared for the four sample fluids: pure water, aqueous solution of sodium dodecyl sulfate (SDS), water-based ZnO nanofluid and water-based ZnO nanofluid in the presence of SDS (0.001 mass fraction). The results showed that the time of hydrate growth decreased and the amount of the storage gas enhanced in the presence of nanoparticles. The literature survey indicated that the nanofluids can be effectively used for improving the performance of several hydrate formation processes. The studies involving some of industrially important nanoparticle systems, such as Al$_2$O$_3$, SiO$_2$, Ag and Cu nanoparticles, on the performance of CO$_2$–CH$_4$ hydrate formation are not yet been reported in the open literature.

In this study, the effects of Al$_2$O$_3$, SiO$_2$, Ag and Cu nanoparticles on the kinetics of CO$_2$–CH$_4$ hydrate formation process were experimentally studied by measuring the amount of gas consumed and the rate of gas consumption. The use of single apparatus, with exactly the same experimental conditions, allowed us to compare the effectiveness of these different nanoparticles.

2. EXPERIMENTAL PROCEDURE

2.1. Compounds used

The gases used (CO$_2$ and CH$_4$) were provided by the Air-Liquide Company. The CO$_2$–CH$_4$ mixture was prepared to be representative of a gas stream rich in CO$_2$ (75 mol% CH$_4$ and 25 mol% CO$_2$). The water used was distilled through a water purifier (Milli-Q plus185 model), and the total organic carbon rate was estimated to be less than 50%. A suspension of 0.1wt%, 0.2wt % and 0.3wt% of different nanoparticles (Al$_2$O$_3$, SiO$_2$, Cu and Ag) was prepared and tested. Table 1 shows some physio-chemical properties of the used nanoparticles.
Table 1: Physio-chemical properties of nanoparticles used

<table>
<thead>
<tr>
<th>Nanoparticles</th>
<th>Diameter (nm)</th>
<th>Purity (%)</th>
<th>Density (g/cm³)</th>
<th>Specific Surface area (m²/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al₂O₃</td>
<td>&lt;50</td>
<td>&gt;99.0</td>
<td>4.00</td>
<td>&gt;40 (BET)</td>
</tr>
<tr>
<td>SiO₂</td>
<td>10-20</td>
<td>&gt;99.5</td>
<td>2.2-2.60</td>
<td>140-180</td>
</tr>
<tr>
<td>Cu</td>
<td>40-60</td>
<td>&gt;99.9</td>
<td>8.94</td>
<td>~12</td>
</tr>
<tr>
<td>Ag</td>
<td>&lt;150</td>
<td>&gt;99.0</td>
<td>10.49</td>
<td>&gt; 5.4</td>
</tr>
</tbody>
</table>

2.2. Compounds used

The main experiment apparatus (Figure 1) is the reactor in which the hydrates are formed and dissociated. The tank is enclosed in an autoclave equipped with sapphire glass portholes. The portholes enable visual observation of the reactor inside. The autoclave is cooled or heated using a cryostat (RLM LAUDA 6S). The refrigerant mixture (water + ethanol) circulates through a double wall to ensure heat exchange within the autoclave. The liquid introduced into the reactor can be stirred with a stirrer, at a speed of up to 800 RPM. A high pressure pump (PU-1587 of JASCO Corporation) allows the injection of the liquid into the pressurized reactor. A vacuum pump (ALCATEL brand) allows us to purge the reactor of any remnants from previous experiments. A cylindrical bottle is connected to the main reactor by a system of tubes and valves, and acts as the supply unit of the pressurized gas mixture. A sampling system called ROLSI (Rapid On-Line Sampler Injector), developed by Mines Paris Tech, was fixed on the top of the autoclave. The gas phase within the reactor could be analyzed by an in-line gas chromatograph (Varian 450 GC model) which was equipped with a TCD detector.
2.3. Procedure

2.3.1. Nanofluid preparation

A suspension of 0.1 wt%, 0.2 wt% and 0.3 wt% of different nanoparticles (Al$_2$O$_3$, SiO$_2$, Cu and Ag) was prepared. To prepare these nanofluids, the required amounts of nanoparticles were dispersed in 5 ml of demineralized water and stirred for 2 hours (to ensure that there was no caking). The resultant solution was then further diluted with distilled water to reach a volume of 400 ml. This dilution procedure is well explained in the experimental procedure section.

2.3.2. Formation of CO$_2$-CH$_4$ mixed hydrates

2.3.2.1 Experimental procedure with pure water

At the beginning of each test, the reactor was evacuated using the vacuum pump in order to eliminate all traces of the previous experiment, and then pressurized to 4 MPa ($P_{exp}$) to initiate a new experiment. The gas phase was stirred and cooled to a temperature of 1°C ($T_{exp}$). Once this temperature was reached, the stirring was stopped. At this time, an analysis of the gas phase by the in-line GC was carried out to check the initial composition. At time $t=t_a$, 400 ml of pure water was injected into the reactor using a HPLC pump (the selected
volume can be changed in accordance with the ability to see the gas-liquid interface through the reactor portholes). The stirring was started again, and set at 350 RPM (which corresponds to the speed limit to avoid vortex and bubbles formation). From this moment onwards, the gas phase composition was analyzed every two hours by the GC. As soon as the water was injected, the temperature and pressure increased simultaneously. The reactor was cooled again and brought down to the temperature of the experiment (1°C), which was kept constant until equilibrium was reached. The evolution of the key parameters (pressure, temperature) of this experiment is shown in Figure 2a.

2.3.2.2 Experimental procedure with nanofluids

Nanofluids of different nanoparticles concentrations of (0.1, 0.2 and 0.3 wt%) were previously prepared by the dispersion of precise amounts (0.4 to 1.2g) of them in 5 ml of distilled water. Here, the procedure followed for 0.1 wt% of Al₂O₃ is described, and is consistent with the procedure followed for other nanofluids as well. At time t=0, 5 ml of the first sample of Al₂O₃ nanofluid (0.4g/5 ml of pure water) was introduced into the reactor, which was then pressurized with the gas mixture to a pressure of 4 MPa (P_{exp}). The gas phase was stirred and cooled to a temperature of 1°C (T_{exp}). At this time, stirring was stopped, and an analysis of the gas phase was performed using the GC to check the initial composition of the mixture within the reactor. At time t=ta, 395 ml of pure water was injected into the reactor using the HPLC pump. The stirring was started again at a speed of 350 RPM. A different procedure is followed here, when compared to the case of pure water in order to avoid difficulties in cleaning nanoparticles from the HPLC pump. From this point on, the procedure was identical to that of the previous case with pure water. Figure 2b shows the evolution of key parameters being measured (P, T) during this experiment (nanofluid of 0.3 %wt SiO₂/water).
**Figure 2a:** Evolution of pressure and temperature for the gas hydrate formation in pure water.

**Figure 2b:** Evolution of Pressure and Temperature for the gas hydrate formation with nanofluid (0.3%wt SiO$_2$)
As is shown in Figures 2a and 2b, there is simultaneous increase in the pressure and temperature during the first stage of the experiment, which corresponds to the injection of water into the reactor, then, there is an initial sharp decrease in the pressure, which corresponds to the dissolution of gas in the aqueous phase until it is supersaturated with respect to the equilibrium conditions. After that, the pressure in the reactor decreases due to the hydrate growth. As can be seen in these two figures, a slight increase in the temperature of the system is observed exactly at the beginning of the hydrate growth stage. This increase in temperature is owing to the exothermic nature of the hydrate formation. The pressure is rapidly decreases in the presence of nanoparticles compared to the case of water and the equilibrium pressure is achieved after 80 hours with nanoparticles; however, in water this pressure is achieved after 170 hours.

3. RESULTS AND DISCUSSION

3.1.Dissolution

To understand the kinetics of dissolution, the amount of gas consumed between the times \( t_0 \) (the time at which the gas is injected) and \( t_a \) (the time at which the crystallization begins) may be tracked. The rate constant of the dissolution of gas in water \( K_{t\alpha} \), now becomes an important factor in the study of this process. Using the following consumption rate equation (1), the value of \( K_{t\alpha} \) is calculated:

\[
r = K_{t\alpha}(C - C_s)
\]  

(1)

Where:

\( r \): Rate of gas consumption and dissolution (mol.s\(^{-1}\))

\( K_t \): Liquid-side mass transfer coefficient (m.s\(^{-1}\))

\( \alpha \): Specific interfacial area based on the liquid volume (m\(^{-1}\))

\( C \): Gas concentration at the gas-liquid interface (mol/m\(^3\))
Gas concentration in the liquid (mol/m$^3$)

Experimentally, the total amount of gas consumed ($A_t$) during dissolution at the time ($t$) can be calculated from the pressure, $P(t)$, and the reactor temperature, $T(t)$, obtained from the data acquisition device connected to the reactor. This amount of gas consumed can by expressed by the following equation:

$$A_t = \frac{V_G}{R} \left( \frac{P_i}{Z_i T_i} - \frac{P(t)}{Z(t) T(t)} \right) \quad (2)$$

$P$, $V_G$ and $T$ are the pressure, volume, and temperature of the gas, respectively. $R$ is the gas universal constant and $Z$ is the compressibility factor.

In this expression, the index $i$ refers to the initial time at the beginning of the experiment, just before the liquid is injected into the reactor. At this point in time, the amount of gas consumed is assumed to be zero. The compressibility factors, $Z_i$ and $Z(t)$, can be determined from the Peng-Robinson equation (Mohammadi et al., 2014; Najibi et al., 2015).

At any time $t$:

$$A_f - A_t = (C_{ext} - C_t)V_G \quad (3)$$

Where:

$A_f$: The cumulative consumption of the gas mixture at saturation

$A_t$: The cumulative consumption of the gas mixture at time $t$

$C_{ext}$: The gas concentration at the gas-liquid interface

$C_t$: The gas concentration in the liquid at time $t$

The rate of consumption is written as:

$$\frac{dA_t}{dt} = K_t \alpha (C_{ext} - C_t)V_G = K_t \alpha (A_f - A_t) \quad (4)$$

By integrating this equation between the time $t_0$, the start time of stirring, and any point of time $t$ the following relationship may be obtained:
\[ \ln \frac{A_f}{A_f - A_t} = K_i(t - t_0) \quad (5) \]

At time \( t_0 \), the gas consumption may be assumed to be zero.

### 3.1.1. Evolution of pressure during dissolution of the gas mixture

Figures 3a through 3d represent the evolution of the pressure ratio \( P/P_s \) (\( P_s \) corresponds to the pressure immediately after the injection of water) as a function of time during the dissolution phase of the gas mixture in pure water and in nanofluids of \( \text{Al}_2\text{O}_3 \), \( \text{SiO}_2 \), Cu and Ag, respectively.

**Figure 3a:** Pressure evolution during dissolution (\( \text{Al}_2\text{O}_3 \) nanofluid)
**Figure 3b:** Pressure evolution during dissolution (SiO$_2$ nanofluid)

**Figure 3c:** Pressure evolution during dissolution (Cu nanofluid)
Figure 3d: Pressure evolution during dissolution (Ag nanofluid)

The $P/P_s$ ratio is observed to rapidly decrease in the presence of SiO$_2$ nanoparticles as compared to pure water, particularly for the concentration of 0.2wt% and 0.3 wt%. Different decreases of this ratio are noted in presence of Cu and Al$_2$O$_3$ nanoparticles, however, no change is observed with Ag nanoparticles.

3.1.2. Gas consumption during dissolution

Figures 4a trough 4d show the consumption of gas versus time during the dissolution of the gas mixture in pure water and in nanofluids Al$_2$O$_3$, SiO$_2$, Cu and Ag, respectively. Figure 4e shows the gas consumption improvement during the dissolution compared to the case of pure water.
Figure 4a: Gas consumption during dissolution (Al₂O₃ nanofluid)

Figure 4b: Gas consumption during dissolution (SiO₂ nanofluid)
Figure 4c: Gas consumption during dissolution (Cu nanofluid)

Figure 4d: Gas consumption during dissolution (Ag nanofluid)
Figure 4e: Gas consumption improvement compared to the case of pure water during dissolution.

To explain the obtained results during this stage, it is interesting to note that:

First, regarding the tested nanoparticles type, the SiO$_2$ particles had the highest specific surface, while Ag particles had the lowest surface area (Table 1), the decreasing in the pressure during the dissolution is due to the dissolution of gas in the aqueous phase (mass transfer) until it is supersaturated and this dissolution depends on the gas-liquid interface area, it can be assumed that may be the presence of nanoparticles with high specific area (eg. SiO$_2$) in the hydrate formation system increases the interfacial area, which enhances the mass transfer (gas consumption).

Second, regarding the concentration of nanoparticles used, it can be hypothesized that there may be bounds within which nanoparticle concentration must fall for these solutions to be effective during the gas dissolution stage. These bounds would be of two kinds: a lower
bound, below which the presence of these particles has a negligible impact on increasing of gas-liquid interfacial area; and an upper bound, above which other factors (such as agglomeration and layer “barrier” formation at gas-liquid interface) may interfere with any benefits these nanoparticles may provide. These bounding concentrations vary from one nanoparticle to the next, but, as can be seen from those that were tested, a high specific surface area seems to lead to strong positive promotion at certain concentration levels. This lower bound is demonstrated by nanoparticles of silica, where very low effect is seen at concentration of 0.1 wt%, while the upper bound can be observed in the case of Cu, where, at concentration 0.2 wt% the kinetics revert to those of pure water.

3.1.3. Calculation of the dissolution rate constant $K_a$ for different nanofluids

Figures 5a through 5d show the evolution of the dissolution rate constant of the gas mixture, $K_a$, in pure water and in nanofluids of Al$_2$O$_3$, SiO$_2$, Cu and Ag, respectively. Figures 6a through 6d look at the initial linear part of this curve for the same nanofluids.

![Graph showing dissolution rate constant for different nanofluids](image)

**Figure 5a:** Calculation of Dissolution rate constant (Al$_2$O$_3$ nanofluid)
**Figure 6a**: Calculation of Dissolution rate constant-Linear part of curve (Al$_2$O$_3$ nanofluid)

**Figure 5b**: Calculation of Dissolution rate constant (SiO$_2$ nanofluid)
Figure 6b: Calculation of Dissolution rate constant-Linear part of curve (SiO₂ nanofluid)

Figure 5c: Calculation of Dissolution rate constant (Cu nanofluid)
**Figure 6c:** Calculation of Dissolution rate constant-Linear part of curve (Cu nanofluid)

**Figure 5d:** Calculation of Dissolution rate constant (Ag nanofluid)
Figure 6d: Calculation of Dissolution rate constant-Linear part of curve (Ag nanofluid)

Initially, these trends are perfectly linear, but gradually deviate from linearity as saturation approaches. The alignment of the experimental points indicates that dissolution kinetics is not disturbed by the use of nanofluids. The kinetic model remains valid, and the determination of constants of the gas dissolution rate (\(K_a\)) can be done by evaluating the slope of the linear part of the curve.

The values of \(K_a\) reported in Table 2 are for the gas mixture in the presence of different nanofluids with a stirring rate of 350 RPM.

The transfer constant is observed to increase slightly for nanofluids of Al\(_2\)O\(_3\) and SiO\(_2\), particularly for SiO\(_2\) at 0.3%. However, \(K_a\) seems to decrease in the case of Cu and Ag nanofluids. From the literature (Olle et al.2006), an enhancement in \(K_a\) may be attributed to either an increase in mass transfer coefficient (\(K_f\)) or an increase in an interfacial area (\(a\)) or both. In this work, it seems like the presence of nanoparticles with high specific surface area is improving the interface area (\(a\)) and consequently improve the \(K_{i\alpha}\).
Table 2: Values of transfer constant $K_1a$ for different nanofluids

<table>
<thead>
<tr>
<th>Nanofluid</th>
<th>$K_1a$ $[s^{-1}]$ $(10^{-4})$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure Water</td>
<td>5.20</td>
</tr>
<tr>
<td>$Al_2O_3/H_2O$</td>
<td>0.1wt% 5.60</td>
</tr>
<tr>
<td></td>
<td>0.2wt% 5.90</td>
</tr>
<tr>
<td></td>
<td>0.3wt% 4.80</td>
</tr>
<tr>
<td>$SiO_2/H_2O$</td>
<td>0.1wt% 5.60</td>
</tr>
<tr>
<td></td>
<td>0.2wt% 5.80</td>
</tr>
<tr>
<td></td>
<td>0.3wt% 12.00</td>
</tr>
<tr>
<td>$Cu/H_2O$</td>
<td>0.1wt% 4.00</td>
</tr>
<tr>
<td></td>
<td>0.2wt% 3.90</td>
</tr>
<tr>
<td></td>
<td>0.3wt% 2.00</td>
</tr>
<tr>
<td>$Ag/H_2O$</td>
<td>0.1wt% 4.00</td>
</tr>
<tr>
<td></td>
<td>0.2wt% 4.90</td>
</tr>
</tbody>
</table>

3.2. Crystallization

Crystallization rates are obtained by evaluating the amount of gas consumed as a function of time during the hydrate formation process. Here, the initial time corresponds to the onset of the exothermic peak or the visual detection of the first hydrate crystals. The evolution of pressure in the reactor, combined with the gas composition (obtained via chromatograph measurements), allows the determination of the crystallization rates of each gas.

3.2.1. Evolution of Pressure during hydrate crystallization

Figures 7a through 7d show the evolution of the pressure ratio $P/P_c$ (where $P_c$ is the pressure at the onset of hydrate crystallization) as a function of time during hydrate crystallization in pure water and in nanofluids of $Al_2O_3$, $SiO_2$, $Cu$ and $Ag$, respectively.
Figure 7a: Evolution of Pressure with time during hydrate crystallization (Al$_2$O$_3$ nanofluid)

Figure 7b: Evolution of Pressure with time during hydrate crystallization (SiO$_2$ nanofluid)
Figure 7c: Evolution of Pressure with time during hydrate crystallization (Cu nanofluid)

Figure 7d: Evolution of Pressure with time during hydrate crystallization (Ag nanofluid)
These results show that in the presence of Al₂O₃ nanoparticles, the $P/P_c$ ratio decreases rapidly for concentrations of 0.1 wt% and 0.2 wt%, but not for the concentration of 0.3 wt%. In the case of SiO₂ nanofluids, a very rapid decline in the $P/P_c$ ratio is observed for the concentration of 0.3 wt%, while no significant effects were observed for the other two concentrations. The solutions of Cu nanoparticles in pure water with 0.1 wt%, 0.2 wt% and 0.3 wt% show a considerable influence on the $P/P_c$ ratio at the beginning of the experiment, but it converges to the effectiveness of pure water by the time the experiments end. At the beginning of crystallization, the presence of Ag nanoparticles seems to have a positive effect on the pressure ratio, but this gradually reverses by the end of crystallization.

3.2.2. Moles of gas consumed during hydrate crystallization

Figures 8a through 8d show the moles of gas consumed per unit volume as a function of time during hydrate crystallization with various nanofluids. Figure 8e shows the gas consumption improvement during the crystallization compared to the case of pure water.

![Graph showing gas consumption during crystallization](image)

**Figure 8a:** Gas consumption during crystallization (Al₂O₃ nanofluid)
Figure 8b: Gas consumption during crystallization (SiO$_2$ nanofluid)

Figure 8c: Gas consumption during crystallization (Cu nanofluid)
**Figure 8d:** Gas consumption during crystallization (Ag nanofluid)

**Figure 8e:** Gas consumption improvement compared to the case of pure water during crystallization.
The presence of SiO$_2$ nanoparticles in pure water at concentrations of 0.3 wt% causes a considerable increase in the amount of gas consumed during hydrate crystallization (average gas consumption improvement of 77%); while small change is visible at the other two concentrations. In the case of Al$_2$O$_3$ nanoparticles, there is an important increase in the amount of gas consumed during crystallization at concentrations of 0.2 wt% and 0.3 wt% (average gas consumption improvement of 30% to 65%); however, this does not seem to be the case at a concentration of 0.1 wt%. Cu nanoparticles at 0.1 wt%, 0.2 wt% and 0.3 wt% show a considerable influence on the amount of gas consumed at the beginning of the experiment, but it converges to the effectiveness by the end of this phase. A positive influence is shown by Ag nanoparticles at the beginning of the crystallization process; however, this reverses to a negative one at the later stages of the experiment, in line with our observations on pressure.

These obtained results can be explained as follow:

First, regarding the type of tested nanoparticles, the hydrates formation process is an exothermic nature, which means a considerable amount of heat is released when hydrate crystals are growing in the system, also for the hydrate former gas with low solubility in water, such CO$_2$, the mass transport of the hydrate former to the hydrate surface may dominate the growth process. Based on the demonstrated positive effect of nanoparticles with high specific surface area on mass and heat transfer phenomena in fluids, it can be assumed that the addition of nanoparticles with high specific surface area (eg. SiO$_2$) increased strongly the mass and heat transfer between the gases, liquid, and hydrate phases, which increased the amount of consumed gas and the released heat and consequently the improvement of crystallization.

Second, regarding the concentration of nanoparticles used, it can be assumed that there may be bounds within which nanoparticle concentration must fall for these solutions to
be effective during the crystallization. These bounds would be of two kinds: a lower bound, below which the presence of these particles has a negligible impact on increasing of heat transfer; and an upper bound, above which other factors (such as agglomeration) may interfere with any benefits these nanoparticles may provide. These bounding concentrations vary from one nanoparticle to the next, but, as can be seen from those that were tested, a high specific surface area seems to lead to strong positive promotion at certain concentration levels. This lower bound is demonstrated by nanoparticles of silica, where very low effect is seen at concentration of 0.1 wt%, while the upper bound can be observed in the case of Cu, where, at concentration 0.2 wt% the kinetics revert to those of pure water.

3.2.3. Variation of the gas consumption rate during hydrate crystallization in pure water and in different nanofluids

The evolution of the gas consumption rate as a function of time during the hydrate crystallization process in nanofluids of Al\textsubscript{2}O\textsubscript{3}, SiO\textsubscript{2}, Cu and Ag are shown in Figures 9a through 9d, respectively.

**Figure 9a:** Gas consumption rate during crystallization (Al\textsubscript{2}O\textsubscript{3} nanofluid)
Figure 9b: Gas consumption rate during crystallization (SiO2 nanofluid)

Figure 9c: Gas consumption rate during crystallization (Cu nanofluid)
Figure 9d: Gas consumption rate during crystallization (Ag nanofluid)

The presence of Al$_2$O$_3$ nanoparticles at concentrations of 0.1 wt\% and 0.2 wt\% considerably increases the gas consumption rate during crystallization, while no significant change of this rate appears at a concentration of 0.3 wt\%. The addition of SiO$_2$ nanoparticles also causes a significant increase in the gas consumption rate during crystallization at a concentration of 0.1 wt\%, but little or no effect is observed for the other two concentrations. In the case of Cu nanoparticles with concentrations of 0.1 wt\%, 0.2 wt\% and 0.3 wt\%, there is a significant positive influence on the gas consumption rate during the crystallization at the beginning of the experiment, but this eventually converges with results for pure water. Similarly, at the beginning of the crystallization step, the presence of Ag nanoparticles has a positive effect on the gas consumption rate, but this effect also seems to reach a limiting behavior at the end of the process.

3.2.4. Evolution of the mole fraction of CO$_2$ and CH$_4$ during hydrate crystallization

Figures 10a through 10d show the evolution of the mole fraction of CO$_2$ in the gas phase as a function of time during the hydrate crystallization process in nanofluids of Al$_2$O$_3$,
SiO₂, Cu and Ag, respectively. **Figures 11a through 11d** show the evolution of the mole fraction of CH₄ in the gas phase as a function of time during the hydrate crystallization process for the same nanofluids.

**Figure 10a:** Evolution of mole fraction of CO₂ in the gas phase during crystallization (Al₂O₃ nanofluid)

**Figure 11a:** Evolution of mole fraction of CH₄ in the gas phase during crystallization (Al₂O₃ nanofluid)
Figure 10b: Evolution of mole fraction of CO₂ in the gas phase during crystallization (SiO₂ nanofluid)

Figure 11b: Evolution of mole fraction of CH₄ in the gas phase during crystallization (SiO₂ nanofluid)
Figure 10c: Evolution of mole fraction of CO$_2$ in the gas phase during crystallization (Cu nanofluid)

![Graph of CO$_2$ mole fraction vs time for different Cu nanofluid concentrations.]

Figure 11c: Evolution of mole fraction of CH$_4$ in the gas phase during crystallization (Cu nanofluid)

![Graph of CH$_4$ mole fraction vs time for different Cu nanofluid concentrations.]

**Figure 10d:** Evolution of mole fraction of CO\(_2\) in the gas phase during crystallization (Ag nanofluid)

**Figure 11d:** Evolution of mole fraction of CH\(_4\) in the gas phase during crystallization (Ag nanofluid)
The addition of Al₂O₃ nanoparticles to pure water at a concentration of 0.3 wt% causes a slight increase in the capture selectivity of CO₂, as compared to CH₄ during crystallization, while no significant change in this selectivity appears for concentrations of 0.1 wt% and 0.2 wt%. Similarly, the presence of SiO₂ nanoparticles, a marginal increase in the capture selectivity of CO₂ compared to CH₄ during crystallization at a concentration of 0.3 wt% has been observed. However, no effect is observed for the other two concentrations. In the case of Cu nanoparticles, none of the three concentrations tested significantly impacted the selectivity of CO₂ capture. Finally, the addition of Ag nanoparticles at concentrations of 0.1 wt%, 0.2 wt% and 0.3 wt% negatively impacts the CO₂ capture selectivity as compared to CH₄.

4. CONCLUSION

The effect of Al₂O₃, SiO₂, Ag and Cu nanoparticles on the kinetics of hydrate formation of a gas mixture rich in CO₂ (75 mol% CH₄ and 25 mol% CO₂) was studied. The results obtained show that the presence of these nanoparticles induces mixed effects on the improvement of the kinetics of this process, the degree of which seems to differ from one additive to another, and also with the concentration in nanoparticles being used. During dissolution, it was observed that nanoparticles of SiO₂ had the most positive effect on gas consumption, particularly concentration of 0.3 wt%. At this concentration the average amount of gas consumed was about 45% higher than that in pure. Cu and Al₂O₃ nanoparticles had an intermediate effect with average improvement of 1% to 15%. Ag nanoparticles had no significant effect. It can be assumed that may be the presence of nanoparticles with high specific area (eg. SiO₂) in the hydrate formation system increased the interfacial area, which enhances the mass transfer (gas consumption). During crystallization, it was seen that nano-silica at a concentration of 0.3 wt% had a very important effect on gas consumption with an improvement of 77% compared to the pure water. In the case of alumina, the significant
impact was observed between 0.1 and 0.2 wt%, the gas consumption was increased by 30% to 65%, while Ag nanoparticles had no significant effect during this phase. It can be hypothesized that the addition of nanoparticles with high specific surface area increased strongly the mass and heat transfer between the gases, liquid, and hydrate phases, which increased the amount of consumed gas and the released heat.

5. ACKNOWLEDGMENT

LCMVAR (Laboratory of Materials Chemistry and the Living), University of Batna is gratefully acknowledged for financial support of this work. ENSM Saint-Etienne (France), Sonatrach Upstream Activity, Technologies & Development Division and Indian Institute of Technology Madras are gratefully acknowledged for technical support and advisory.

6. REFERENCES

Ansari, F., Sobhani, A., Salavati-Niasari, Masoud,. 2014. Sol–gel auto-combustion synthesis of PbFe\textsubscript{12}O\textsubscript{19} using maltose as a novel reductant. RSC Advances. 4, 63946-63950


