Gas Hydrate Formation Process for Capture of Carbon Dioxide from Fuel Gas Mixture

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To determine the suitable operating conditions for the hydrate-based CO2 separation process from a fuel gas mixture, the hydrate nucleation and growth kinetics of the simulated fuel gas (39.2 mol % CO2/H2 gas mixture) in the presence of tetra-n-butyl ammonium bromide (TBAB) are investigated. The experiments were conducted at the TBAB concentration range of 0.14–1.00 mol %, the temperature range of 275.15–282.45 K, the driving force range of 1.00–4.50 MPa, the gas/liquid phase ratio range of 0.86–6.47, and the hydrate growth time of 15–120 min. It is found that the addition of TBAB not only shortens the induction time and accelerates the hydrate growth rate, but also enhances CO2 encaged into the hydrate. However, the number of total moles of gas consumed and the number of moles of CO2 transferred into the hydrate slurry phase decrease with the increase of the TBAB concentration when the TBAB concentration is above 0.29 mol %. The induction time reduces, and the number of moles of gas consumed, the hydrate formation rate, and the number of moles of CO2 encaged into hydrate phase increase with the increase of the driving force. However, when the driving force is more than 2.5 MPa, H2 prefers to go into the hydrate phase with the increase of the driving force, as compared to CO2. In addition, the temperature has little effect on the hydrate formation process.

1. Introduction

It is acknowledged that the emission of carbon dioxide from the burning of fossil fuels is the major contributor to the greenhouse gas emission and subsequent global warming and climate change. Particularly, fossil fuel electric power plants are producing about one-third of CO2 emission worldwide.1 A promising approach to provide the near future electricity from fossil fuels for the worldwide increasing energy requirements with near-zero CO2 emissions is integrated gasification combined cycle (IGCC) technology with precombustion CO2 capture.2 An essential element of this approach, in fact, is CO2 separation from the mixture of CO2 and H2. Removal of CO2 from the gas mixture can be achieved by a number of separation techniques including absorption into a liquid solvent, adsorption onto a solid, cryogenic separation, and permeation through membranes. However, these methods have the individual issues of either high corrosion, large energy consumption, high cost, or low capacity, and so forth.3 Hence, it is urgent to develop an efficient and cost-effective capture/separation technology for the CO2 separation.

A novel method for separating CO2 is via gas hydrate crystallization.4–7 The basis for the separation is the selective partition of the CO2 component between the hydrate phase and the gaseous phase. Spencer et al.8 gave an economic assessment that the cost of the hydrate technique for CO2 separation from IGCC power plant is approximately 10 U.S. dollars per ton of CO2, which is much lower than that of other methods. Hence, the hydrate separation technique is promising for separating CO2 from IGCC fuel gas in the power plants.

Linga et al.5 studied the hydrate kinetics of the CO2/H2 mixture in pure water system. They found that the induction time is 9.7 min and the hydrate formation rate is 0.0048 mol/min for the first 5 min for the CO2/H2/H2O system at 273.7 K and 7.5 MPa. Especially, the operating pressure of 7.5 MPa was higher than the outlet pressure of the fuel gas. Hence, there is a continuous interest in using the additives to shorten the induction time, accelerate the hydrate formation rate, and reduce the operating pressure. Some promising additives for the fuel gas mixture are propane (C3H8), cyclopentane (CP), tetrahydrofuran (THF), etc. Zhang et al.5 proposed the hydrate-based CO2 capturing process from the precombustion stream with CP. They found that the processes of the two stages of the hydrate formation and dissociation can significantly enrich CO2 from 40 to 98 mol % at 282 K. Lee et al.5 found that THF of 1.0 mol % is the optimum concentration for the CO2 capture based on the kinetic experiments. On the basis of kinetic experiments coupled with the compositional analysis, Kumar et al.7 proposed a two-stage hydrate/membrane process with an addition of 2.5 mol % C3H8 into the fuel gas mixture. Recently, tetra-n-butyl ammonium bromide (TBAB) has taken a particular attraction as an environmentally friendly additive for the gas separation due to small gas molecules (CH4, CO2) that can be engaged into the dodecahedral cavities (S-cage, 512) of the TBAB hydrate at favorable stability conditions.9 Duc et al.10 presented the thermodynamic data showing that the addition of 0.29 mol % TBAB can substantially decrease the formation pressure of the CO2/N2 hydrate. Moreover, the CO2 selectivity in the hydrate phase is at least 4 times better than that in the gas phase. Fan et al.11 presented the kinetic data showing that the addition of 0.293 mol % TBAB can substantially decrease the formation pressure of the CO2/N2 hydrate. However,
new studies on the kinetic behavior of the hydrate formation for the CO2/H2 gas mixture in the presence of TBAB are reported.

The objective of the present work is to investigate into the effects of the TBAB concentration, the gas/liquid phase volume ratio, the driving force, and the experimental temperature on the separation of CO2 from the CO2/H2 gas mixture based on the kinetic experiments coupled with the compositional analysis and the resistance detecting method.

2. Experimental Section

2.1. Materials. A treated synthesis gas coming out of an IGCC power station consists of approximately 40 mol % CO2/H2 gas mixture at a total pressure of 2.5–5 MPa. Thus, a CO2/H2 gas mixture containing 39.2 mol % CO2 was used in the work to simulate a pretreated fuel gas mixture. Tetra-n-butyl ammonium bromide (TBAB) with 99.9% purity was supplied by Shanghai Sinopharm Chemical Reagent Co., Ltd., China. The deionized water with the resistivity of 18.25 MΩ cm−1 used in the work was produced by an ultrapure water system supplied by Nanjing Ultrapure Water Technology Co., Ltd., China.

2.2. Apparatus. The experimental apparatus used in this work is shown in Figure 1. The inner volume and the maximum working pressure of the high-pressure crystallizer (CR) are 336 cm3 and 25 MPa, respectively. The crystallizer has two circular viewing windows on the front and the back. Mixing of the crystallizer contents is accomplished using a magnetic stir bar (450 r/min) that is magnetically coupled to a set of the rotating magnet, which is driven by an electric motor (Shanghai Meiyinpu instrument, Ltd., China). Two Pt1000 thermoprobes (JM6081) with ±0.1 K accuracy are used to measure the temperatures in the gas phase and in the hydrate slurry phase in the crystallizer, respectively. All pressure measurements are determined with Setra smart pressure transducers (model 552, Boxborough, MA, U.S.) with the pressure range of 0–25 MPa and the accuracy of ±0.02 MPa. To measure the composition of the gas phase and the hydrate slurry phase in the crystallizer, a Wufeng GC522 gas chromatographer (GC) (Shanghai Wufeng Scientific Instrument CO., Ltd., China) is connected online with the crystallizer and automated with a personal computer (PC). The crystallizer (CR) and the supply vessel (SV) are immersed in a temperature-controlled bath containing a 30/70 wt % ethylene glycol/water mixture.

2.3. Procedure. The procedure followed in this work is described in detail elsewhere. Prior to the experiment, the crystallizer is cleaned using the deionized water and allowed to dry. Next, the TBAB aqueous solution prepared at a desired concentration was injected into the high-pressure hydrate crystallizer to a desired volume. Subsequently, the hydrate crystallizer with the solution was flushed with the CO2/H2 mixture gas at least four times to remove any residual air, and then it was filled with the CO2/H2 mixture gas until the desired pressure is reached. Once the temperature was stabilized (typically within 1 min) the stirrer in the crystallizer was started, and the experimental time also began to be recorded. During the experiment, the temperature, the pressure, and the resistance in the system were recorded once every 6 s, and the residual gas was sampled and its composition was analyzed with GC at the 5th, 15th, 35th, 60th, and 90th minute. As the gas in the crystallizer was consumed on account of the hydrate formation, additional gas was supplied, and the pressure in the crystallizer was maintained constant with a proportional integral derivative (PID) controller.

The equilibrium hydrate formation conditions for the CO2/H2/TBAB/H2O mixtures are required for the kinetic experiments and have been determined in our previous work. Each gas uptake experiment was carried out at the individual driving force (Pexp − Peq), which is the difference of the experimental pressure (Pexp) from the equilibrium pressure (Peq). Table 1 provides the list of the experimental conditions.

2.4. Amount of Gas Consumed. The number of moles of the gas (ΔnH2) that has either gone into the water or been consumed from the hydrate formation can be calculated as follows:

\[ \Delta n_{H2} = n_{H2}^{eq} - n_{H2}^{0} = \left( \frac{P}{zRT} \right)_{G,0}^{SV,0} - \left( \frac{P}{zRT} \right)_{SV,t} \]

where z is the compressibility factor calculated by Pitzer’s correlation, subscript t refers to time t, subscript 0 refers to the initial time, subscript G refers to the gas phase in the crystallizer, and subscript SV refers to the gas phase in the supply vessel.

During the kinetic experiment, the composition of the gas phase is determined by GC. At any given time by knowing the composition of the gas mixture in the crystallizer, the number of moles of the individual gas component consumed for the hydrate formation can be calculated by the component mass balance.

\[ \Delta n_{i}^{H2} = n_{i}^{H2} - n_{i}^{H2,0} = V_{G}\left[ \frac{(y_{i}^{H2}P^{H2})}{zRT_{G,0}} - \frac{(y_{i}^{H2}P^{H2})}{zRT_{SV,0}} + \frac{y_{i}^{H2}PV}{zRT_{SV,0}} - \frac{y_{i}^{H2}PV}{zRT_{SV,t}} \right] \]

where superscript t refers to the component of the gas mixture, and y is the mole fraction. It is assumed that the volume changes due to phase transitions are neglected.

3. Results and Discussion

In this work, a total of 18 kinetic experimental runs were carried out at the different conditions including the fresh and memory water (runs 1 and 2), the gas/liquid phase volume ratio (runs 2–5), the experimental temperature (runs 2, 6–8), the TBAB concentration (runs 2, 9–12), and the driving force (runs 2, 13–18) to find the suitable operating condition for CO2.
the system resistance slightly reduces on account of the effect
the pressure of the SV slightly decreases with the slight increase
slightly increases from 278.15 to 278.25 K due to the stirring,
time of the hydrate formation. During the period, the temperature
C to
is the process of extensive hydrate formation, which takes
is because a small amount of the hydrate continues forming
increasing gradually, and the pressure continues reducing slowly
in the increase of the temperature in the system. In this period,
the pressure of the SV reduces quickly with the rapid increase
of the number of the gas consumption on account of the
extensive gas hydrate formation. The phenomena are similar to
those observed by Linga et al. and Lee et al.4,6 In addition, it is
found that the system resistance gradually increases over time
in the process. It is due to the fact that the solid hydrate formed
by water, dissolved CO2, the ions of Br, and tetra-n-butyl
ammonium causes the decreases of the concentrations of the
ions of Br and tetra-n-butyl ammonium in the TBAB solution.
Thus, the decrease of the number of moles of the free ions in
the solution and the increase of the amount of the hydrate slurry
in the process of the hydrate formation result in the reduction
of the electric conductivity in the system, that is, the increase
of the system resistance. In the period from C to D, the
temperature has little change, the system resistance continues
increasing gradually, and the pressure continues reducing slowly
with the slow increase of the moles of the consumed gas. This
is because a small amount of the hydrate continues forming
during the period. The process takes approximately 20 min.
After point D, the temperature, the pressure, the resistance, and
the consumption of the gas all have no change with time. This
illustrates that the hydrate formation has completed at point D.
A similar process can also be observed in other runs.

3.2. Effect of Memory Water. Memory refers to the situation
where water that is used in the experiment has experienced the
hydrate formation.13 Table 1 gives the induction time for the
hydrate formation from the systems with fresh and memory
water is 5.8 min, and it is obviously shorter than that with fresh
water. 13.4 min. This observation agrees with what is described
by Bishnoi and Natarajan.16 Actually, the operation with
memory water is practical and feasible for the industrial
continuous loop. Thus, the following experiments were con-
ducted with memory water.

3.3. Effect of Gas/Liquid Phase Volume Ratio. The gas/
liquid phase volume ratio (Rv) is defined as the ratio of the
system sharply increases from 278.25 to 278.85 K within a short
time, and then gradually decreases to 278.15 K. This period is
a process of the phase transformation of gas and water to solid
hydrate, which creates the heat of formation and results
in the increase of the temperature in the system. In this period,
the pressure of the SV reduces quickly with the rapid increase
of the number of the gas consumption on account of the
effective gas hydrate formation. The phenomena are similar to
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found that the system resistance gradually increases over time
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by water, dissolved CO2, the ions of Br, and tetra-n-butyl
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ions of Br and tetra-n-butyl ammonium in the TBAB solution.
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Table 1. Experimental Conditions along with Measured Induction Time and Hydrate Formation Rates for Different Systems

<table>
<thead>
<tr>
<th>items</th>
<th>F/M</th>
<th>x, mol %</th>
<th>Rv, kPa</th>
<th>T, °C</th>
<th>DF, kPa</th>
<th>time, min</th>
<th>time1/2, min</th>
<th>n, mol</th>
<th>Rv, kPa/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>run 1</td>
<td>F</td>
<td>0.29</td>
<td>0.86</td>
<td>278.15</td>
<td>2.50</td>
<td>90</td>
<td>13.4</td>
<td>3.82 × 10⁻³</td>
<td>7.26 × 10⁻³</td>
</tr>
<tr>
<td>run 2</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>278.15</td>
<td>2.50</td>
<td>90</td>
<td>5.8</td>
<td>2.02 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 3</td>
<td>M</td>
<td>0.29</td>
<td>1.49</td>
<td>278.15</td>
<td>2.50</td>
<td>90</td>
<td>5.9</td>
<td>1.98 × 10⁻³</td>
<td>7.23 × 10⁻³</td>
</tr>
<tr>
<td>run 6</td>
<td>M</td>
<td>0.29</td>
<td>1.49</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>5.9</td>
<td>1.98 × 10⁻³</td>
<td>7.23 × 10⁻³</td>
</tr>
<tr>
<td>run 7</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>279.55</td>
<td>2.50</td>
<td>90</td>
<td>6.1</td>
<td>1.99 × 10⁻³</td>
<td>7.28 × 10⁻³</td>
</tr>
<tr>
<td>run 8</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>279.55</td>
<td>2.50</td>
<td>90</td>
<td>6.2</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 9</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>279.55</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 10</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 11</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 12</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 13</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 14</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 15</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 16</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 17</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
<tr>
<td>run 18</td>
<td>M</td>
<td>0.29</td>
<td>0.86</td>
<td>277.05</td>
<td>2.50</td>
<td>90</td>
<td>6.3</td>
<td>2.06 × 10⁻³</td>
<td>7.32 × 10⁻³</td>
</tr>
</tbody>
</table>

*Fresh water. †Memory water. ‡TBAB concentration in aqueous solution. §Gas/liquid phase volume ratio. ¶Experimental temperature. †Driving force (Pexp − Pvap). ‡Reacted time. §Induction time. ¶Moles of gas consumed at nucleation point (induction time). †Rate of hydrate growth (gas consumption rate for the first 5 min after nucleation).
volume of the gas phase \( (V_G) \) to the volume of the liquid phase \( (V_L) \) in the crystallizer. Figure 3 gives the comparison of the gas uptake curves for the hydrate formation with the gas/liquid phase volume ratio of 0.86–6.47 in the presence of 0.29 mol % TBAB at 278.15 K and the driving force of 2.50 MPa, as shown in runs 2 to 5. As seen from the Figure 3, the number of the total moles of the gas consumed during hydrate formation increases as \( R_v \) reduces from 6.47 to 0.86. For example, the number of moles of gas consumed at the end of the hydrate formation is up to 0.1044 mol at \( R_v = 0.86 \), while it is only 0.0497 mol at \( R_v = 6.47 \). This interesting phenomenon can be explained as follows: because the inter volume of the crystallizer \( (V_C) \) is constant, \( R_v = (V_C - V_L)/V_L = V_G/V_L - 1 \) is only a function of the volume of the liquid phase \( (V_L) \) in the crystallizer, and the value of \( R_v \) decreases with the increase of the amount of the TBAB solution. That is, in this work, the liquid phase volume increases from 45 to 180 mL as the gas/liquid phase volume ratio decreases from 6.47 to 0.86. The bigger liquid phase volume means more moles of TBAB existing in the solution for the hydrate formation. Thus, at the given condition, the more moles of TBAB in the solution results in the more TBAB hydrom formed with more enclosed gas. This is why the moles of gas consumed increase as the \( (R_v) \) decreases. Therefore, in the following experiments, \( R_v = 0.86 \) corresponding to the liquid volume of 180 mL was chosen to obtain the highest gas consumption resulting in the highest amount of the formed hydrate. In addition, it is found that the gas/liquid phase volume ratio has no effect on the induction time, as shown in Figure 3.

3.4. Effect of Experimental Temperature. Figure 4 shows the gas uptake measurements for hydrate formation from the system with 0.29 mol % TBAB at the driving force of 2.50 MPa and the different temperatures, as shown in runs 2, 6–8. The results show that the value of the gas uptake and the gas consumption rate at the fixed time and the induction time have no significant change with the change of the temperature. The above phenomenon illustrates that the temperature change has little effect on the kinetic behavior of the hydrate formation with the given driving force.

3.5. Effect of TBAB Concentration. As shown in runs 2, 9–12, for the systems with the TBAB concentration of 0.14–1.00 mol %, the experiments were carried out at the temperatures whose equilibrium hydrate formation pressure is 0.50 MPa with the above different TBAB concentrations. The data of the equilibrium hydrate formation condition have been measured in our previous work. Thus, all the experimental pressures are fixed at 3.0 MPa to obtain the driving force of 2.50 MPa. The experimental data are summarized in Table 1 (2, 9–12) and plotted in Figure 5. It can be seen from Figure 5 that the hydrate forms quickly with the TBAB concentration from 0.21 to 1.00 mol % except for the TBAB concentration of 0.14 mol %, for which the induction time is 9.5 min, and the induction time reduces with the increase of the TBAB concentration. However, it is not the higher TBAB concentration that causes the more gas consumed. The number of total moles of gas consumed increases with the increase of TBAB concentration in the lower concentration range. However, it decreases with increasing the TBAB concentration when the TBAB concentration is more than 0.29 mol %. For example, the number of total moles of gas consumed with the TBAB concentrations of 0.14, 0.21, and 0.29 mol % is 0.0897, 0.0988, and 0.1047...
mol, respectively, which exhibits an increase trend, whereas, with the TBAB concentrations of 0.50 and 1.00 mol %, the number of total moles of the gas consumed is only 0.0949 and 0.0894 mol, respectively, which shows a decrease trend. It may be due to the fact that the gas hydrate sharply forms substantially and agglomerates at the gas/liquid interface when TBAB concentration is relatively high. Extensive hydrate formation and crystal agglomeration results in the accumulation of crystals as stagnant pockets at the gas/water interface, which prevents more gas from coming into contact with the water. A similar phenomenon can be found elsewhere. The more agglomeration results in the smaller amount of the gas into the solution for the hydrate formation.

The residual gas compositions in the systems with 0.14–1.00 mol % TBAB at driving force of 2.50 MPa are given in Figure 6. As seen in Figure 6, the CO₂ composition in the residual gas phase in the system with the given TBAB concentration decreases with time from 0 to 60 min, and then slightly increases with time from 60 to 90 min. For example, the CO₂ composition in the residual gas phase in the system with 0.29 mol % TBAB substantially decreases from 39.2 to 18.36 mol % with time from 0 to 60 min, and then slightly rises to 18.45 mol % at the 90th minute. On the other hand, in the systems with the different TBAB concentrations, the CO₂ composition in the residual gas during the hydrate growth decreases as the TBAB concentration increases from 0.14 to 0.29 mol % at a certain time. However, it is not the higher TBAB concentration that causes the lower CO₂ composition in the residual gas. On the contrary, when the TBAB concentration is more than 0.29 mol %, the CO₂ composition in the residual gas is higher than that with 0.29 mol % TBAB at the given time after 15 min and increases with the increase of the TBAB concentration. It is because, as mentioned above, the gas hydrate sharply forms substantially and agglomerates at the gas/liquid interface when the TBAB concentration is relatively high. The agglomeration of the hydrate at the gas/liquid interface hinders the further mass transfer of CO₂ into the aqueous solution. Furthermore, this causes the increase of the CO₂ composition in the residual gas phase with the high concentration of TBAB.

### 3.6. Effect of Driving Force

Figure 7 shows the comparison of the gas uptake curves for the hydrate formation from the system with 0.29 mol % TBAB at 278.15 K and different driving forces. As seen from Figure 7, the induction time decreases with the increase of the driving force. For example, the induction time was 43 min with 1.0 MPa driving force, while it is only 3.0 min with 4.5 MPa driving force. In addition, the gas consumption rate and the total moles of gas consumed increase with increasing the driving force. This is attributed to the fact that, on one hand, the higher driving force makes more gas going into the aqueous solution and furthermore causes more gas hydrate formed, resulting in more gas consumed; on the other hand, the increase of the amount of the gas going into the solution correlates with the enhancement of the gas hydrate growth rate, which also means the enhancement of the gas consumption rate. Figure 8 shows the changes of the CO₂ compositions in the residual gas.
phase versus time under the same experimental conditions with Figure 7. As seen in Figure 8, the CO_2 composition in the residual gas phase decreases substantially with time at the given driving force except for 1.0 MPa. It is for this reason that CO_2 in the gas phase reduce with time. However, it does not mean that the higher driving force causes the lower CO_2 composition in the residual gas phase. For example, the CO_2 composition in the residual gas phase decreases as the driving force increases from 1.0 to 2.5 MPa at the certain time. On the contrary, the CO_2 content in the residual gas phase increases as the driving force increases from 3.0 to 4.5 MPa, which can be attributed to the fact that H_2 competes with CO_2 for hydrate cage (5^{12}) occupancy at the higher driving force. The observed phenomenon is in agreement with the report by Linga et al.4

4. Conclusions

The hydrate nucleation and growth kinetics of the CO_2/H_2/ TBAB hydrate systems are investigated at the different conditions. The total moles of gas consumed increase with the gas/liquid phase volume ratio, which has no effect on the induction time. The driving force can shorten the induction time and promote the hydrate formation. The induction time reduces, and the total moles of the consumed gas corresponding to the amount of the hydrate formation, the hydrate formation pressure of CO_2 is much lower than that of H_2. Furthermore, the process makes the CO_2 content in the residual gas phase reduce with time. However, it does not mean that the higher driving force causes the lower CO_2 composition in the residual gas phase. For example, the CO_2 composition in the residual gas phase decreases as the driving force increases from 1.0 to 2.5 MPa at the certain time. On the contrary, the CO_2 content in the residual gas phase increases as the driving force increases from 3.0 to 4.5 MPa, which can be attributed to the fact that H_2 competes with CO_2 for hydrate cage (5^{12}) occupancy at the higher driving force. The observed phenomenon is in agreement with the report by Linga et al.4

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