Effects of Composite Accelerators on the Formation of Carbon Dioxide Hydrates

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ABSTRACT: To improve the rate of formation of carbon dioxide hydrates, tetra-n-butylammonium bromide (TBAB) was compounded with different concentrations of sodium dodecyl sulfate (SDS) and nanographite, and the effects of these mixtures on carbon dioxide hydrate formation were studied. The addition of TBAB alone, as well as mixtures of TBAB and SDS or nanographite, shortened the induced nucleation time, and the induction times of the TBAB−2.5 g/L nanographite and TBAB−0.24 g/L SDS systems were the shortest and longest, respectively. Further, on mixing TBAB and SDS, the induced nucleation time first increased and then decreased with the increase in the SDS concentration. When TBAB and nanographite were mixed together, the induced nucleation time first decreased, then increased, and again decreased with the increase in the nanographite concentration. In addition, the hydrate formation rate and conversion were highest for the TBAB−0.48 g/L SDS system and lowest for the TBAB−0.06 g/L SDS system; in the first 35 min, from the end of gas charging, the TBAB−10 g/L nanographite and TBAB−5 g/L nanographite systems yielded the highest and lowest hydrate formation rates and conversions, respectively. For the composite systems, obvious effects were observed in the initial stages of reaction, but the effects varied over the course of the reaction. Overall, the use of different accelerators resulted in little differences in the total production, conversion, and formation rate of carbon dioxide hydrates over the course of the reaction.

1. INTRODUCTION

Natural gas hydrate technology has been widely used in gas storage and transportation, seawater desalination, CO₂ storage, cold-storage technology, and transportation; however, owing to the slow hydrate formation rate and low gas storage capacity, its large-scale industrial utilization has been limited. Therefore, researchers have focused on increasing the rate of hydrate generation. Physical and chemical methods can be used to increase this rate. Physical methods involve expanding the gas−liquid contact area and enhancing the heat- and mass-transfer capacity by bubbling, stirring, or spraying. In contrast, in the chemical method, the formation of hydrates is enhanced by adding accelerators, which allow the use of milder reaction conditions and increase the amount of dissolved gas. Concerning energy consumption and cost, physical methods are less favorable than chemical methods, so chemical methods have proven to be popular. Common accelerators include thermodynamic accelerators, kinetic accelerators, nanofluids, ionic accelerators, biosurfactants, and others.

Among them, surfactants and kinetic accelerators have attracted the most attention because they show the best enhancement of hydrate formation. For example, Ye et al. found that the addition of tetra-n-butylammonium bromide (TBAB) in mass fractions of 0.1 and 0.05 accelerates the rate of formation of carbon dioxide hydrates, and the addition of 0.05 wt % TBAB was optimal. Babu et al. tested the effect of the addition of 0.3, 1, 1.5, 2, and 3 mol % TBAB and found that the 1 mol % TBAB solution resulted in the highest rate of carbon dioxide hydrate formation. Further, the induction time when using the 0.3 mol % TBAB solution was the longest, and the overall gas consumption was the largest because this solution had high gas solubility before nucleation. Nguyen et al. found that the use of a solution containing 0.125% TBAB reduced the carbon dioxide hydrate formation rate, thus acting as an inhibitor. Therefore, accelerators can show different effects on the hydrate formation rate at different concentrations. Further, for a single accelerator, there is an optimal concentration, but this concentration changes with temperature, pressure, and other factors.

Received: December 3, 2021
Accepted: March 30, 2022
Published: April 29, 2022
In detail, Babu et al.,27 Zhou et al.,34 and Lin et al.35 found that on carbon dioxide hydrate formation at 272.65 K and 3.3 MPa. Using the optimal TBAB concentration (vide infra), we studied the use of SDS and nanographite alone; therefore, in this study, enhanced the formation of carbon dioxide hydrates compared to shown that mixtures of SDS, TBAB, and nanographite can optimal combinations and concentrations. Previous research has effects of different accelerator combinations. Therefore, it is necessary to study the effects of different accelerator types and concentrations on carbon dioxide hydrate formation and identify the optimal combinations and concentrations. Previous research has shown that mixtures of SDS, TBAB, and nanographite can enhance the formation of carbon dioxide hydrates compared to the use of SDS and nanographite alone; therefore, in this study, using the optimal TBAB concentration (vide infra), we studied the effects of different concentrations of SDS and nanographite on carbon dioxide hydrate formation at 272.65 K and 3.3 MPa. In detail, Babu et al.,27 Zhou et al.,34 and Lin et al.35 found that the optimum concentrations of TBAB, nanographite, and SDS for carbon dioxide hydrate formation to be 0.288, S, and 0.24 g/L respectively, so TBAB was combined with 0.06, 0.12, 0.24, 0.36, or 0.48 g/L SDS or 1.25, 2.5, 5, 7.5, and 10 g/L nanographite. Thus, we identified the most suitable accelerator system for enhancing carbon dioxide hydrate formation. The findings of this study will aid the industrialization of carbon dioxide hydrate formation and the rapid capture of carbon dioxide, thus enabling the large-scale utilization of hydrates.

2. EXPERIMENTAL SECTION

2.1. Materials. CO$_2$ was provided by Lanzhou Zhongkekeate Co. Ltd. (purity: 99.99%). Deionized water was prepared in our laboratory and had a conductivity of 18.25 mΩ cm. SDS was provided by Chuang Baiyou Biological Reagent Consumables (purity: 99.9%). Nanographite was provided by Jishengya Nano Technology Co. Ltd. (10,000 mesh; purity: 99.9%). TBAB was provided by Shanghai McLean Biochemical Technology Co. Ltd. and had a purity of 99.9%.

2.2. Apparatus. The experimental system used in this experiment is shown in Figure 1 (yellow represents the circuitry

![Figure 1. Carbon dioxide hydrate formation apparatus.](https://doi.org/10.1021/acsomega.1c06834)
the same height each time to ensure that the same sample volume was used. The reactor and sensor were then installed, and the temperature of the circulating cold bath was adjusted to 272.65 K. Simultaneously, vacuum was applied to the solution for 10 min. After reaching the set temperature, the precooled gas was charged into the reactor to 3.3 MPa to initiate hydrate formation. The hydrate formation process was considered to have been completed when the pressure and temperature in the reactor did not vary for 48 h after the formation of the carbon dioxide hydrate.

### 2.4. Data Processing

The amount, rate, and conversion of carbon dioxide hydrate were considered during analysis and experiment. The calculation process is as follows.

#### 2.4.1. Amount of Generated Carbon Dioxide Hydrate

The amount of carbon dioxide hydrate formed was determined at different periods of the experiment: during the gas charging process and from the end of gas charging to the end of the experiment. The amount of hydrate generated during gas charging, \( n_o \), is the difference between that produced over the entire experiment (beginning to the end of the experiment) and that generated from the end of gas charging to the end of the experiment, as given by eq 1.

\[
n_t = n_d - n_e
\]

Here, \( n_o \), \( n_p \), and \( n_e \) are the amounts of hydrate generated during gas charging, over the entire experiment, and from the end of gas charging to the end of the experiment (mol), respectively.

The percentage of volume generated during gas charging with respect to the whole generated volume is given by eq 2.

\[
n_{10} = \frac{n_t}{n_d} \times 100\%
\]

Here, \( n_{10} \) is the percentage of volume generated during gas charging with respect to the whole generated volume.

Using the experimental temperature and pressure and the equation for hydrate formation, the amount of hydrate in the reactor from the end of gas charging to the end of the experiment, \( n_{i1} \), was calculated, as shown in eq 7.

\[
n_{1i}H_2O + CO_2 = CO_2\cdot n_{i}H_2O
\]

Here, \( n_i \) is the hydration number. The amount of TBAB used was \( 8.93 \times 10^{-4} \) mol, and the amount of carbon dioxide hydrate generated by the final TBAB system is 0.3457 mol; thus, the TBAB accounts for 2.58\% of the final amount of hydrate. Because of this low value, the semi-cage hydrate generated by TBAB and pure ice powder was ignored, and the hydration number was assumed to be 5.75\% under ideal conditions.

\[
n_{o} = \frac{P_{o}V}{Z_{o}RT_{o}}
\]

Here, \( P_{o} \) and \( T_{o} \) are the pressure (MPa) and temperature (K), respectively, in the reactor at the end of inflation, \( V \) is the volume (L) occupied by the gas in the reactor at the end of inflation, \( R \) is the universal gas constant (8.314 J/(mol K)), \( Z_{o} \) is the gas compression factor at the end of inflation, which was obtained by iteration of the Peng–Robinson (PR) equation calculated by Li et al.\(^{27}\) and Smith.\(^{38}\) \( n_{o} \) is the amount of carbon dioxide hydrate in the reactor at the end of inflation.

The amount of carbon dioxide hydrate in the reactor from the end of inflation to the end of the formation experiment, \( n_{i0} \), was calculated using eq 5.

\[
n_{i0} = \frac{P_{i}V}{Z_{i}RT_{i}}
\]

Here, \( P_{i} \) and \( T_{i} \) are the pressure (MPa) and temperature (K), respectively, at time \( t \) from the end of inflation to the end of the formation experiment, and \( Z_{i} \) is the gas compression factor at time \( t \) from the point when gas charging stopped to the end of the experiment; this was also obtained using the PR equation reported by Li et al.\(^{27}\) and Smith.\(^{38}\) Finally, \( n_{i} \) is the amount of carbon dioxide hydrate in the reactor at time \( t \) from when gas charging stopped to the end of the generation experiment.

Therefore, the amount of carbon dioxide hydrate generated at any time from the end of charging to the end of the experiment, \( \Delta n \), is given by eq 6.

\[
\Delta n = n_{o} - n_{i} = \frac{P_{o}V}{Z_{o}RT_{o}} - \frac{P_{i}V}{Z_{i}RT_{i}}
\]

The generation amount from the end of inflations to the end of the experiment, \( n_{p} \), was calculated, as shown in eq 7.

\[
n_{pi} = n_{o} - n_{i} = \frac{P_{o}V}{Z_{o}RT_{o}} - \frac{P_{i}V}{Z_{i}RT_{i}}
\]

Here, \( P_{1} \) and \( T_{1} \) are the pressure (MPa) and temperature (K) at the end of the generation experiment, respectively, and \( Z_{1} \) is the gas compression factor at the end of the generation experiment, which was obtained using the PR equation as proposed by Li et al.\(^{27}\) and Smith.\(^{38}\) \( n_{i} \) is the amount of carbon dioxide hydrate in the reactor at the end of the generation experiment.

#### 2.4.2. Rate of CO2 Hydrate Formation.

The average rate of carbon dioxide hydrate formation was assessed during the gas charging process and from the beginning of gas charging to the end of the formation experiment. The formation rate during the charging process, \( v \), is the ratio of the amount of carbon dioxide hydrate generated during charging from the beginning of induced nucleation to the end of charging, as given by eq 8.

\[
v = \frac{n_{i}t}{t_{i}}
\]

Here, \( v \) has units of mol/min and \( t_{i} \) is the time from the beginning of induced nucleation time to the end of inflation (min).

The formation rate\(^{38}\) from the end of gas charging to the end of the experiment was calculated every 5 min. Using eq 8, the average formation rate of carbon dioxide hydrate every 5 min (\( v \)) can be calculated.

\[
v = \frac{n_{i} - n_{i+5}}{5} \quad (i = i + 5, i = 0)
\]

Here, \( n_{i} \) is the amount of carbon dioxide hydrate in the reactor from the end of gas charging to \( i \) min and to the end of the experiment and \( n_{i+5} \) is the amount of carbon dioxide hydrate in the reactor at \( i + 5 \) min.

#### 2.4.3. Ice Conversion Rate

The ice conversion rate is the ratio of ice consumed before and after the reaction to the amount of ice in the reactor during the experiment. In this study, we investigated the conversion rate during the gas charging process and from the end of gas charging to the end of the experiment. The conversion rate, \( \eta_{c} \), during gas charging is the difference between the conversion rate from the beginning to the end of the experiment, \( \eta_{p} \), and that from the beginning of gas charging to the end of the experiment, \( \eta_{o} \), as given by eq 13.
\[ \eta_t = \eta_d - \eta_g \tag{10} \]

The total conversion rate \( \eta_t \) is calculated based on the measured total production, that is, conversion rate \( \eta_g \) from the beginning to the end of the experiment and is given by eq 11.

\[ \eta_g = \frac{5.75n_d}{n_i} \tag{11} \]

The conversion rate at any time from the stop of inflation to the end of the generation experiment, \( \Delta \eta_g \), is given by eq 12.

\[ \Delta \eta_g = \frac{5.75\Delta n}{n_i} \tag{12} \]

In addition, eq 13 provides the conversion rate from the end of gas charging to the end of the experiment

\[ \eta_d = \frac{5.75n_k}{n_i} \tag{13} \]

Here, as above, \( n_i \) is the amount of ice in the reactor and \( \eta_d \) is the ice conversion rate.

3. RESULTS AND DISCUSSION

3.1. Determination of the Induction Period and Hydrate Formation Phase. To enable the comparison of the effects of SDS and nanographite on hydrate formation, the formation of carbon dioxide hydrate using a pure ice powder—nitrogen-cooled system and a TBAB-only system was studied. The experiments involving different accelerators are labeled 1–12, as shown in Table 2. Except for the pure ice—nitrogen gas system, the trends in the temperature and pressure of all experiments were similar, so we discuss one representative case only. After the formation of carbon dioxide hydrate, the temperature reached the set temperature and the pressure stabilized; the overall hydrate formation time was 24 h. The temperature and pressure changes during carbon dioxide hydrate formation in the TBAB and pure ice—nitrogen gas systems are shown in Figure 2. As indicated in the figure, there are three main stages: the induced nucleation period (marked 0–A in the figure), rapid hydrate formation, and an equilibrium region.

From point 0–A in Figure 2, the temperature of the TBAB accelerator system first rises, then decreases slightly, and then rises again. In the same time period, the pure ice—nitrogen system shows a rapid increase, followed by a slight decrease in temperature. Based on these results, no significant hydrate formation occurs in this time period, which corresponds to the induced nucleation period. For the TBAB system, the point at which the temperature rises rapidly and intersects with the nitrogen temperature curve and the corresponding pressure decreases indicates the start of rapid hydrate formation. Thus, this is the hydrate-formation stage. Figure 3 shows the corresponding results for pure ice powder—nitrogen, pure ice powder—TBAB, and pure ice powder—TBAB—1.25 g/L nanographite systems. The induced nucleation periods for these last two systems are marked 0–A1 and 0–A2, respectively. From point A1, massive hydrate formation starts in these systems. In Figure 2, points A–D indicate the rapid hydrate formation stage. Point C indicates the end of gas charging. In particular, from points C to D, the pressure drops sharply, indicating the formation of a large number of hydrates. In contrast, from point D to the end of the experiment, the pressure decreases slowly until it plateaus, and the reactor temperature then gradually stabilizes at 272.65 K, indicating that hydrate formation is complete.

3.2. Effect of Different Accelerator Systems in the Induction Period. Figures 4 and 5 show the induced nucleation times for hydrate formation for the different systems. As shown, the induced nucleation time for the compound systems is shorter than that of the pure ice powder system, which is consistent with the findings of Zhang et al.29 and Zhou et al.31 who also found that the addition of SDS and nanographite shortens the induction period for carbon dioxide formation.

![Figure 2. Temperature and pressure during the formation of carbon dioxide hydrate in the TBAB and pure ice systems.](https://doi.org/10.1021/acsomega.1c06834)
hydrate formation. Crucially, there are significant differences between the induction times of the SDS and nanographite systems.

As shown in Figure 4, the induced nucleation time first increases and then decreases with increase in the SDS concentration, and the samples containing 0.06 and 0.24 g/L SDS have the shortest and longest induction times, respectively. Excluding the 0.24 g/L SDS system, all induced nucleation times of the TBAB−SDS system are shorter than that of TBAB alone, indicating that the compound accelerator enhances nucleation. Further, although higher SDS concentrations shorten the induction period, there is an optimal value.

As shown in Figure 5, for the nanographite system, the induction time decreases, then increases, and then decreases again with increase in the nanographite concentration, and the induction times of the 2.5 and 7.25 g/L nanographite systems were the shortest and longest, respectively. The TBAB−nanographite system also showed shorter nucleation times than that of TBAB alone.

In conclusion, for the compound systems, the 2.5 g/L nanographite and 0.24 g/L SDS system yielded the shortest and longest, respectively, induction times.

### 3.3. Effects of Different Accelerant Systems on the Production of CO₂ Hydrate

The amount of hydrate formation mainly includes the amount of gas charging process and the amount of gas hydrate formation from the stop of gas charging to the end of the formation experiment. The amount of hydrate generated during gas charging is the total amount minus that generated once gas charging stops, as shown in Table 2. The temperature and pressure at which all the experiments were conducted were 273.65 K and 3.3 MPa, respectively. During gas charging, the maximum amount of hydrate was formed for the TBAB−0.48 g/L SDS system (0.1174 mol, which accounts for 22.67% of the total amount) and the minimum amount was formed for the TBAB−0.12 g/L SDS system (0.0304 mol, which accounts for 8.21% of the total amount).

From the end of the addition of gas to the end of the experiment, the amounts of hydrate formed for cases 1−12 (Table 2) were 0.3435, 0.3457, 0.3381, 0.3397, 0.3411, 0.3359, 0.3352, 0.3358, 0.3338, 0.4048, 0.3381, and 0.3409 mol, respectively. In all cases, there was no significant difference in the amount of hydrate formed from the end of gas charging to the end of the experiments because the temperature, pressure, and ice powder quality were identical, and only the accelerators were different. Dicharry et al. also found that there is little difference in the amount of hydrate formed when adding different accelerators. However, the effect of this experiment is obvious in the first 35 min of inflation stop, mainly studying the formation of the first 35 min.

The changes in the amount of hydrate formed during the course of the experiments are shown in Figures 6 and 7 for the TBAB−SDS and TBAB−nanographite systems, respectively. In these figures, 0 indicates the end of gas charging. For all systems, hydrate formation increased rapidly in the first 5−10 min. Subsequently, hydrate formation slowed and eventually plateaued. Thus, in the first 5 min, the formation rate was very low, but, with increase in time, the formation rate increased. In
the later stages, the carbon dioxide consumption decreased and, thus, the formation of hydrate slowed. The average amounts of hydrate produced for the TBAB−0.48 g/L SDS and TBAB−2.5 g/L nanographite composite systems were the highest in the first 5 min after gas charging stopped. However, with an increase in time, the rate of hydrate generation decreased below those of the other systems; this is because carbon dioxide is consumed rapidly during gas charging, so a large amount of hydrate is formed, and the generation rate between 5 and 10 min is lower than those of the other systems. Over the whole course of the experiment (0−35 min), the formation rate of the TBAB-containing system was greater than that of the pure ice powder because TBAB enhances hydrate formation.40

As shown in Figure 6, at 0−5 min, the maximum (0.0006 mol/g) and minimum (0.0001 mol/g) amounts of hydrate were formed for the TBAB−0.48 g/L SDS and TBAB−0.12 g/L SDS systems, respectively, and there was an optimal SDS concentration. After 10 min, the maximum (0.0033 mol/g) and minimum (0.0030 mol/g) amounts of hydrate were formed for the TBAB−0.06 g/L SDS and TBAB−0.36 g/L SDS systems, respectively. Therefore, the concentration of the TBAB−SDS accelerator had a distinct effect on hydrate formation, and this effect varied over the course of the reaction.

From 0−35 min, hydrate formation in the TBAB-only system was greater than those of the TBAB−SDS systems, and the TBAB−0.06 g/L SDS and TBAB−0.48 g/L SDS systems produced the most and the least, respectively, amount of hydrate. However, both of these values are greater than that of the pure ice powder system. Therefore, the addition of SDS increased the amount of hydrate formation probably because SDS reduces the surface tension and increases the solubility of the hydrate.41,42 However, there was no simple linear, positive correlation between the SDS concentration and hydrate formation.

As shown in Figure 7, at 0−5 min, the maximum (0.0006 mol/g) and minimum (0.0001 mol/g) amounts of hydrate were formed for the TBAB−2.5 g/L nanographite and TBAB−10 g/L nanographite systems, respectively, clearly showing that the concentration of nanographite has a significant effect on hydrate formation. As for the SDS system, the amounts of hydrate formed in the nanographite systems were greater than that formed in the pure ice powder system. At 0−35 min, the maximum amount of hydrate was formed for the TBAB−10 g/L nanographite system (0.0038 mol/g). The reason for the enhanced hydrate formation is that nanographite provides many nucleation sites, thus increasing the hydrate nucleation rate, as well as enhanced heat and mass transfer, thus promoting carbon dioxide hydrate formation.43 The minimum hydrate formation was obtained for the TBAB−5 g/L nanographite system (0.0036 mol/g), suggesting the inhibition of hydrate formation beyond the optimal nanographite concentration.

3.4. Influence of Different Accelerator Systems on the Formation Rate of CO2 Hydrate. The rates of hydrate formation for the different systems are shown in Figures 8 and 9.

In these figures, 0 indicates the end of gas charging, and the values indicated at 0 on the x-axis are the average formation rates from the beginning to the end of gas charging, that is, the average hydrate generation rate during gas charging. During gas charging, the maximum (0.019 mol/min) and minimum (0.005 mol/min) formation rates were obtained for the TBAB systems containing 0.48 and 0.12 g/L SDS, respectively, indicating again that there is an optimal SDS concentration. After gas charging, the average formation rate first increased and then decreased with an increase in the reaction time. At 0−5 min, the maximum hydrate formation rates were obtained for the TBAB systems containing 0.48 g/L SDS and 2.5 g/L nanographite. For the other systems, the formation rate...
Mech et al. found that the hydrate formation rate in a TBAB–reaction in systems containing TBAB and SDS. For example, the critical micelle concentration of SDS, its ability to reduce surface tension and increases hydrate solubility, thus increasing the hydrate nucleation rate, and resulting in a low hydrate formation rate. At 10 g/L nanographite systems, the maximum (0.053 mol/min) and minimum (0.0115 mol/min) hydrate formation rates were obtained for the TBAB–nanographite systems, respectively, but the rates were all greater than that of the pure ice powder system, and there was an optimal TBAB–nanographite concentration. Importantly, the relationship between the concentration and formation rate was not linear, and the effect of the concentration varied with the course of the reaction. Mekala et al. found that the initial rate of carbon dioxide hydrate formation in a brine system was greater than that in a pure water system; however, after the initial stages of the reaction, the formation rate in the pure water system was greater than that of the brine system.

At 0–35 min, the maximum and minimum formation rates were obtained for the TBAB–10 g/L nanographite and TBAB–5 g/L nanographite systems, respectively. In addition, only the TBAB–10 g/L nanographite system had a higher rate of hydrate formation than TBAB alone because of the synergistic interaction of the nanographite and TBAB, which improved the hydrate nucleation rate, making this the optimal nanographite concentration.

### 3.5. Effects of Different Accelerant Systems on Conversion Rate

The rates of conversion of ice to CO₂ hydrate during gas charging are shown in Table 2. For the TBAB–SDS systems, the maximum (21.51%) and minimum (3.58%) conversions were obtained for the TBAB–0.48 g/L SDS and TBAB–0.12 g/L SDS systems, respectively. For the TBAB–nanographite systems, the maximum (10.36%) and minimum (7.22%) conversions were obtained for the TBAB–2.5 g/L nanographite and TBAB–10 g/L nanographite systems, respectively. Among all the complex systems, TBAB–0.48 g/L SDS and TBAB–0.12 g/L SDS had the highest and lowest conversions, respectively. The overall conversion rates for cases 1–12 in Table 2 are 50.94, 51.59, 48.31, 43.58, 51.18, 44.32, 60.98, 47.66, 49.67, 57.79, 49.71, and 47.36%, respectively.

Figures 10 and 11 show the conversion rates for all systems from the end of gas charging to the end of the experiment. In these figures, 0 indicates the end of gas charging, the conversion rate begins to decrease, hindering hydrate formation, and resulting in a low hydrate formation rate. At 10–35 min, the maximum and minimum hydrate formation rates were obtained for the TBAB–0.06 g/L SDS and TBAB–0.48 g/L SDS systems, respectively, but the formation rate of the pure TBAB system was greater than those of the complex systems. Other researchers have also reported differences in the hydrate formation rate over the course of reaction in systems containing TBAB and SDS. For example, Mech et al. found that the hydrate formation rate in a TBAB–600 ppm SDS system was faster than that of TBAB alone in the initial 3 h, but, after 3 h, the rate was significantly slower.

As shown in Figure 9, at 0–5 min, the formation rate of all TBAB and nanographite composite systems is higher than that of the pure ice powder. The reason is that nanographite increases the nucleation rate of hydrate by providing more nucleation points for TBAB molecules. The maximum (0.053 mol/min) and minimum (0.0115 mol/min) hydrate formation rates were obtained for the TBAB–2.5 g/L nanographite and TBAB–10 g/L nanographite systems, respectively, but the rates were all greater than that of the pure ice powder system, and there was an optimal TBAB–nanographite concentration. Importantly, the relationship between the concentration and formation rate was not linear, and the effect of the concentration varied with the course of the reaction. Mekala et al. found that the initial rate of carbon dioxide hydrate formation in a brine system was greater than that in a pure water system; however, after the initial stages of the reaction, the formation rate in the pure water system was greater than that of the brine system.

At 0–35 min, the maximum and minimum formation rates were obtained for the TBAB–10 g/L nanographite and TBAB–5 g/L nanographite systems, respectively. In addition, only the TBAB–10 g/L nanographite system had a higher rate of hydrate formation than TBAB alone because of the synergistic interaction of the nanographite and TBAB, which improved the hydrate nucleation rate, making this the optimal nanographite concentration.

### Conversion Rate

Figure 10. Conversion rates for the TBAB–SDS system at 3.3 MPa and 273.65 K.

Figure 9. Rate of formation of carbon dioxide hydrate for the TBAB–nanographite system at 3.3 MPa and 273.65 K.
pure ice, but the use of TBAB and nanographite reduces this.

The following conclusions can be drawn.

The conversion rates from the end of gas charging to the end of the experiment for cases 1–12 in Table 2 are 40.45, 40.71, 39.81, 40, 40.16, 39.55, 39.47, 39.54, 39.3, 47.66, 39.81, and 40.14%, respectively. Except for the TBAB system, the other composite systems yielded lower conversions than the pure ice powder system, and the conversion of the TBAB–5 g/L nanographite system was the lowest. Further, the conversion varied little from the end of gas charging to the end of the experiment, which is consistent with the results of Lin and Ando. Crucially, the addition of TBAB alone increases the conversion of the hydrate compared to that with pure ice, but the use of TBAB and nanographite reduces this effect. Yu et al. found that nanographite slightly inhibits the formation of carbon dioxide hydrates and affects phase equilibrium. Nashed et al. reported that nanofluids do not significantly promote or inhibit hydrates, and, thus, nanomaterials are mainly used as kinetic accelerators to promote hydrate formation.

4. CONCLUSIONS

The effects of TBAB, SDS, and nanographite accelerators on the formation of carbon dioxide hydrates were studied. The effects of these compounds on hydrate formation were assessed based on the induced nucleation time, speed of hydrate formation, amount of hydrate generated, and conversion of ice to carbon dioxide hydrate. The following conclusions can be drawn.

(1) TBAB alone promoted the formation of carbon dioxide hydrates in the first 35 min after gas charging. The amounts of hydrate produced, formation rate, and ice conversion of the TBAB-only system are greater than those of the pure ice powder system, whereas the induced nucleation time is shorter than that of the pure ice powder system.

(2) The addition of TBAB with SDS or nanographite reduces the induction (nucleation) period. For the TBAB–SDS system, with an increase in the SDS concentration, the induced nucleation time first increased and then decreased, and the TBAB–0.06 g/L SDS and TBAB–0.24 g/L SDS systems yielded the shortest and longest, respectively, induction periods. For the TBAB–nanographite system, the induction time first decreased, then increased, and then decreased again with an increase in the nanographite concentration. The induced nucleation times of the TBAB–2.5 g/L nanographite and TBAB–7.25 g/L nanographite systems are the shortest and the longest, respectively. However, the induced nucleation time with TBAB alone is longer than that with TBAB–nanographite.

(3) The composite accelerators showed few differences in total hydrate production, formation rate, and conversion from the beginning of gas charging to the end of the experiment. Nevertheless, concerning these factors, the 0.48 g/L SDS and TBAB–0.12 g/L SDS systems showed the best and worst performance, respectively. The hydrate formation was the highest in the first 35 min after gas charging stopped, but these effects vary over the course of hydrate formation. In particular, the effects of the accelerators were not obvious after the end of gas charging because hydrate formation occurred at a low rate.

(4) In the first 35 min, after the end of gas charging, the 0.06 g/L SDS system produced the highest amount of hydrate and had the highest generation rate and ice conversion, whereas the 0.12 g/L SDS yielded the lowest generation rate, ice conversion, and generation rate. For the TBAB–nanographite system, these performance metrics for the 10 g/L nanographite and 5 g/L nanographite systems were the best and worst, respectively.

(5) Therefore, to generate hydrates rapidly in large amounts, TBAB is the optimal single accelerator and TBAB and 10 g/L nanographite are the best combined accelerator.

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Funding
This work was supported by the National Natural Science Foundation of China (no. 41661103), Open Fund of National Key Laboratory of Frozen Soil Engineering, Chinese Academy of Sciences (no. SKLFSE201406), and National Key R & D projects (no. 2017YFC0307303).

Notes
The authors declare no competing financial interest.

ACKNOWLEDGMENTS
Thanks to Zhaohui Zhang for helping with these experiments.

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