



Cite this: RSC Adv., 2017, 7, 53563

Investigation of kinetics of tetrabutylammonium chloride (TBAC) + CH₄ semiclathrate hydrate formation

Lingli Shi ^{abc} and Deqing Liang ^{*abc}

For potential application in advanced gas storage at moderate temperatures, a systematic study on tetrabutylammonium chloride (TBAC) + CH₄ semiclathrate hydrate formation kinetics was conducted using isobaric kinetics measurements to evaluate the effects of pressure (6.0, 3.0 MPa), temperature (278 K, subcooling degree of 6 K), and salt concentration (0.10, 0.20, 0.30, 0.34, 0.45 mass fraction). The results revealed that the systems showed shorter induction time, higher normalized gas consumption and higher rapid growth rate under a higher supersaturation environment, represented by higher pressure or lower temperature. Besides, the effect of salt concentration was complicated. With the increase of salt concentration, the total gas consumption was almost the same while the normalized gas consumption decreased greatly, indicating that the amount of CH₄ trapped in the hydrate unit greatly decreased and the system with low salt concentration was a good choice for advanced gas storage. In addition, a Raman device was employed to reveal the structural properties. The spectra showed that the (TBAC + CH₄) semiclathrate hydrates were formed with hexagonal structure or tetragonal structure under different salt concentrations, which were different from the structures of pure TBAC hydrate. It was assumed that at low salt concentrations the addition of CH₄ induced the formation of hexagonal structure since it had three 5¹² cages per TBAC which was higher than that of the tetragonal structure.

Received 25th September 2017

Accepted 15th November 2017

DOI: 10.1039/c7ra10595a

rsc.li/rsc-advances

1. Introduction

Clathrate hydrate is one kind of nonstoichiometric crystalline inclusion compound. It comprises a water lattice held together through hydrogen bonding with guest molecules of gases (such as CH₄, CO₂, or N₂) or some volatile liquids (such as tetrahydrofuran, acetone).¹ Generally, the structures of simple clathrate hydrates are mainly categorized into three structures, namely structure I with space group *Pm3n*, structure II with space group *Fd3m*, and structure H with space group *P6/mmm*, which are formed depending on the size, shape and nature of the accommodated guest species. In cases where the guest molecules are large ionic molecules, such as tetrabutylammonium salts (TBAX, X = F/Cl/Br) and tetrabutylphosphonium salts, the guest cations are engaged in the four cage-fused cage composed by tetrakaidecahedron cages (T, 5¹²6²) and penta-kaidecahedron cages (P, 5¹²6³), and the anions can replace the water molecules in the hydrate cage.^{2–5} The dodecahedron cages (D, 5¹²) are vacant to trap guest gas molecules. Thus this kind of

ionic clathrate hydrate, known as semiclathrate hydrate (SCH), shows many special features, such as cohost inclusion and improved thermal stability, due to the ionic interaction between host and guest molecules.⁶ Recently, SCH has been studied as a more promising medium as compared with ordinary clathrate hydrate for many engineering practices, such as gas separation/storage/transportation,^{7–20} seawater desalination,²¹ and cold energy storage.^{22–25}

SCHs of TBAX (X = F/Cl/Br) were first founded by Fowler *et al.*²⁶ in 1940 and structurally determined by Jeffery²⁷ in 1969. They were reported to be stable at temperatures above 273 K and the most common structures of SCHs were hexagonal structure-I (HS-I), tetragonal structure-I (TS-I) and cubic superstructure-I (CSS-I) with unit cell formula of 2P·2T·3D·40H₂O, 4P·16T·10D·172H₂O, and 48T·16D·368H₂O, respectively.⁴ Rodionova *et al.*³ reported that according to different salt concentrations TBAB, TBAC and TBAF SCHs were formed on the basis of HS-I/TS-I, TS-I, and TS-I/CSS-I, respectively. As compared with some other known thermodynamic promoters, such as tetrahydrofuran (THF), TBAX (X = F/Cl/Br) were more attractive because of their non-volatile property, which could ensure the gas phase uncontaminated and the SCH forming process reversible.

In recent years, many researchers have reported that TBAX (X = F/Cl/Br) could remarkably enhance the stability of gas hydrate by greatly lowering the equilibrium pressure and increasing the equilibrium temperature.^{28–35} While regarding the formation

^aCAS Key Laboratory of Gas Hydrate, Guangzhou Institute of Energy Conversion, Chinese Academy of Sciences, Guangzhou 510640, China

^bGuangdong Provincial Key Laboratory of New and Renewable Energy Research and Development, Guangzhou 510640, China

^cGuangzhou Center for Gas Hydrate Research, Chinese Academy of Sciences, Guangzhou 510640, China. E-mail: liangdq@ms.giec.ac.cn; Fax: +86 2087057669



kinetics which were more important for industrial applications, only some limited experiments have been carried out. Among TBAX ($X = F/Cl/Br$), TBAB was most investigated. In 2013, Roosta *et al.*³⁶ reported that TBAB could increase the CH_4 hydrate formation rate. In 2014, Zhang *et al.*¹⁴ studied the formation kinetics of (TBAB + CO_2) SCHs at mass fraction = (0.05 and 0.10), and the results revealed that the moles of formed SCHs and the formation rates both increased with the increase of subcooling degree. In 2016, Verrent *et al.*³⁷ investigated the hydrate growth of (TBAB + CO_2) at 0.40 salt mass fraction. However, as compared to TBAB, TBAC was reported to give a higher pressure reduction for N_2/CO_2 hydrate formation and a similar pressure reduction for CH_4 hydrate formation.³⁸ To our best knowledge, most of studies related with TBAC were focused on the thermodynamic properties. While considering the structural properties, only a limited data were reported in the open literature.^{4,39–41} Kim *et al.* presented the PXRD patterns of (TBAC + CO_2) SCHs and (TBAC + CO_2 + N_2) SCHs.^{39,40} Oshima *et al.*⁴¹ studied crystallographic properties of (TBAB + TBAC) SCHs through PXRD measurement, and the result demonstrated that the double SCHs were formed with structure of tetragonal one ($P4/mmm$ or $P4/m$) or orthorhombic one ($Pmma$) at different salt concentrations. However, these achievements of TBAC did not involve the formation kinetics which were important for hydrate-based applications. This literature review indicated an imperative need for studying the kinetics of TBAC SCHs with varying mass fractions to understand and develop uses of hydrates, including designing effective processes.

In this work, we studied the formation kinetics of double (TBAC + CH_4) SCHs to determine the effects of TBAC concentration, pressure and temperature on the hydrate nucleation, hydrate growth and gas storage capacity. In addition, the Raman spectroscopy was employed to analyze the hydrate structure and identify the molecular behavior.

2. Experimental section

2.1 Experimental materials

TBAC (0.98 mass fraction, Tokyo Chemical Industry Co., Ltd.), and CH_4 (0.999 mole fraction, Guangzhou Shiyuan Gases Co.,

Ltd.) were purchased and used without further treatment. Deionized water made in the laboratory, after careful degassing, and TBAC were weighed on an electronic balance with the uncertainty of ± 0.01 g to prepare desired TBAC aqueous solution with w (TBAC mass fraction) = (0.10, 0.20, 0.30, 0.34, and 0.45).

2.2 Experimental apparatus

A schematic diagram of the experimental apparatus used for the measurements of hydrate formation kinetics was shown in Fig. 1. The apparatus mainly consisted of a hydrate formation reactor with a stirrer driven by a DC motor, a glycol–water bath, a pressure reducing valve (ER5000SI-1, Tescom), a gas flow meter (EL-FLOW Digital, Bronkhorst), and a data acquisition system (DAQ). The reactor, the core of the apparatus, was made of stainless steel with inner volume of approximately 307 cm³. The glycol–water bath, equipped with two see-through quartz windows (16 mm × 80 mm each), allowed lighting inside the reactor and making video documentation during hydrate formation process. The pressure reducing valve and the gas flow meter were used to maintain a constant pressure with a precision of $\pm 0.5\%$ relative and record the consumed gas, respectively. The temperature and pressure of the system were measured by a platinum resistance with an uncertainty of ± 0.1 K and a pressure transducer with an uncertainty of ± 0.024 MPa, respectively. All the data of temperature, pressure, gas consumption, and process video were collected through the DAQ and saved at preset sampling intervals on a computer.

2.3 Experimental procedures

2.3.1 Measurements of hydrate formation kinetics. The measurement of hydrate formation kinetics was carried out with an isobaric method which was reported in ref. 42. First of all, the reactor was thoroughly washed with deionized water and the aqueous solution twice, respectively. Then, the vacuum reactor was charged with 100 mL of aqueous solution and cooled to the experimental temperature. After the temperature was kept constant for about 0.5 h, the reactor was pressurized to the experimental pressure (6.0, or 3.0 MPa) with CH_4 under the

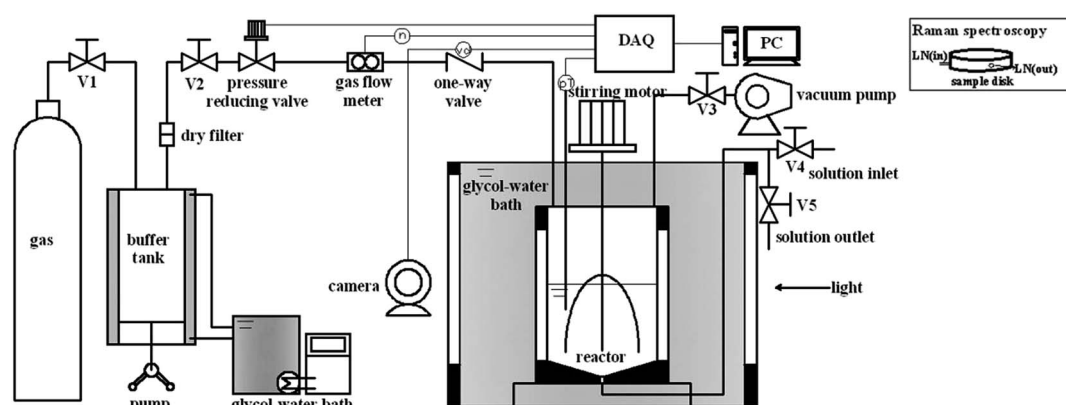


Fig. 1 Schematic diagram of the experimental setup.



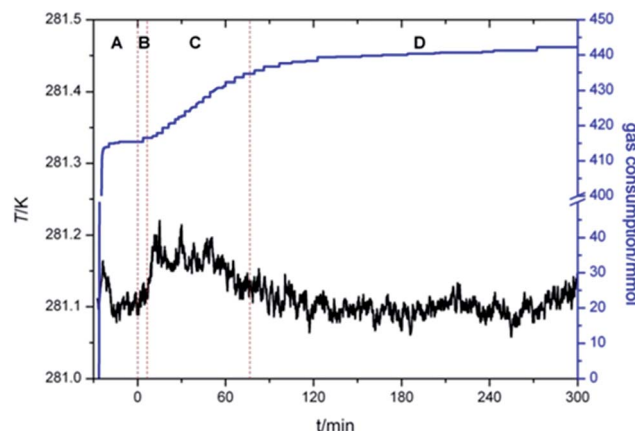


Fig. 2 Temperature and gas consumption versus time in the (TBAC + CH₄) SCh formation process at $w = 0.10$, $P = 3.0$ MPa, and $\Delta T = 6$ K.

control of the pressure reducing valve. Once the temperature increase caused by injected CH₄ was removed, the stirrer was switched on and set to 200 rpm. This value was chosen because the stirring rate was relatively low to observe the hydrate formation process and to compare the effects of temperature, pressure and salt concentration on hydrate kinetics. The end of the hydrate formation could be checked through the stabilized temperature and gas consumption. As shown in Fig. 2, section A was the process in which CH₄ was charged, and sections (B, C and D) were the hydrate formation process, which could be divided into nucleation part, rapid growth part, and slow growth part. All the experiments were repeated thrice for each system with different w , temperatures and pressures to ensure that the experiment results were reliable and reproducible.

2.3.2 Raman measurement. The double (TBAC + CH₄) SChs samples were prepared at P (pressure) = 6.0 MPa and ΔT (subcooling degree) = 6 K. To prevent the dissociation of hydrate shell, the samples were preserved in liquid nitrogen when transported or prepared for microstructural measurement in our laboratory. Raman measurement was performed at 203 K with a confocal Raman spectrometer (LabRAM HR, Horiba) equipped with a multichannel air-cooled CCD detector. An laser source (Torus, Laser Quantum Ltd.) operating at 532 nm was used with a power of 50 mW. The silicon crystal

standard of 520.7 cm⁻¹ was employed to calibrate the subtractive spectrograph. The sample was placed in the sample disk which could keep the sample at 203 K with the help of liquid nitrogen (LN).

3. Results and discussion

3.1 Formation kinetics of (TBAC + CH₄) hydrate

To fully understand the effect of TBAC on CH₄ hydrate formation kinetics, the experiment was conducted with w varying from (0.10 to 0.45) at P = (6.0 and 3.0) MPa under different temperature conditions ($\Delta T = 6$ K, or $T = 278.15$ K). Table 1 and Fig. 3 showed the experimental conditions as well as equilibrium data of double (TBAC + CH₄) SChs.⁴³⁻⁴⁵ It could be seen that the experimental temperatures were higher than the equilibrium temperatures of pure CH₄ hydrate at the same pressure. Hence it could provide a more economical method for hydrated based gas storage. The formation process could be generally divided into three sections: nucleation process, rapid growth process and slow growth process. Thus the investigated parameters included induction time (IT), normalized gas consumption (NGC), hydrate rapid growth time (RGT) and hydrate rapid growth rate (RGR). It was well-known that systems with a higher ΔT and higher P , indicating a higher driving force, were expected to show a better kinetic performance.

Fig. 4 presented the IT of hydrate nucleation process for systems at various w , P and T . The IT was caused by the fact that a supersaturation environment was needed for the hydrate formation. The supersaturation environment was dependent on the pressure and temperature differences between the equilibrium conditions and the operation conditions. In the present study, IT was calculated as the time elapsed from the beginning of the stirring until a significant temperature increase was detected in the system, which could also be confirmed with the assistant of visual observation through the see-through quartz windows. As shown in Fig. 4, under the same ΔT or T , systems with higher P showed shorter IT. And under the same P , systems with lower T also showed shorter IT. While considering the effect of TBAC, IT did not show a regular trend with the increase of w . The stochastic property of IT performed strongly in systems with $w = 0.20$. It should be noted that in systems of (B,

Table 1 Summary of experimental conditions conducted for TBAC + CH₄ + H₂O systems^a

Code	w	P /MPa	T /K	T_{eq} /K	Code	w	P /MPa	T /K	T_{eq} /K
A1	0.10	6.0	283.90	289.9	C3	0.30	3.0	284.35	290.35
A2	0.10	6.0	278.15	289.9	C4	0.30	3.0	278.15	290.35
A3	0.10	3.0	281.10	287.1	D1	0.34	6.0	287.20	293.2
A4	0.10	3.0	278.15	287.1	D2	0.34	6.0	278.15	293.2
B1	0.20	6.0	286.50	292.5	D3	0.34	3.0	284.60	290.6
B2	0.20	6.0	278.15	292.5	D4	0.34	3.0	278.15	290.6
B3	0.20	3.0	283.70	289.7	E1	0.45	6.0	286.30	292.3
B4	0.20	3.0	278.15	289.7	E2	0.45	6.0	278.15	292.3
C1	0.30	6.0	286.90	292.9	E3	0.45	3.0	283.65	289.65
C2	0.30	6.0	278.15	292.9	E4	0.45	3.0	278.15	289.65

^a T_{eq} was the equilibrium data reported in the open literature.



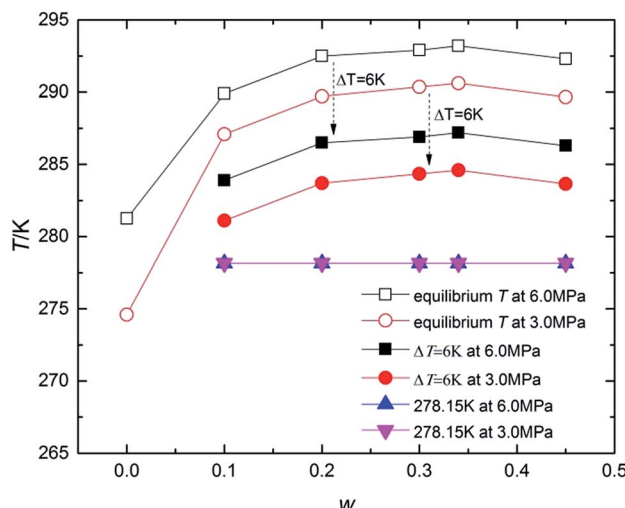


Fig. 3 Kinetic experimental conditions represented on the T - w curve along with reported equilibrium conditions.

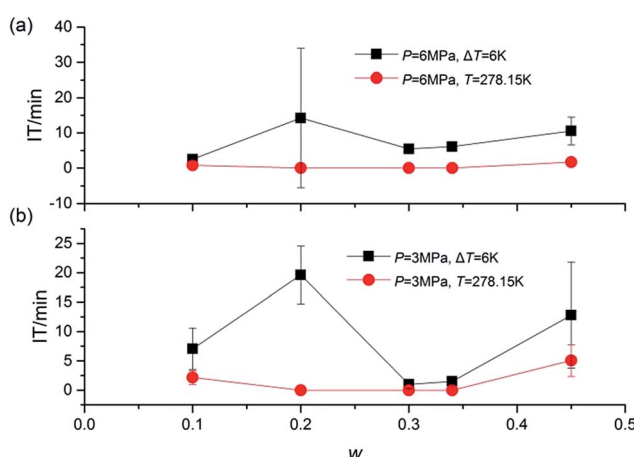


Fig. 4 Induction time (IT) of TBAC + CH_4 hydrate nucleation process at various w with different conditions: (a) $P = 6$ MPa; (b) $P = 3$ MPa. The error bars indicated the standard deviations of each system.

B4, C2, C4, D2 and D4), the hydrate formation had started before the start of stirring, then the IT was set to be 0 in these systems. The readings of stirring were shown only on the digital screen and not collected through DAQ, thus the curve could not be provided. In the future study, the readings would be added in the experimental apparatus. The obvious drawback was the fact that the formation of SCHs seriously hindered the start of stirring. Then it prevented the CH_4 molecules from going into the test fluid and forming double (TBAC + CH_4) SCHs. Thus the prior formation of SCH and unable start of stirring led to the reduction in methane capture. From this viewpoint, the systems with rather long or short IT were not the good choice for industrial applications, since the long IT would cause financial loss and the extremely short IT might lead to the unable start of stirring and the decreased gas storage amount.

Total gas consumption (TGC) was the parameter of hydrate growth process (shown as section C and D in Fig. 2) and

collected with the gas flow meter. As shown in Fig. 5, despite of systems in which stirring was blocked at first, systems with same P showed similar TGC, and it had little difference with the change of w . Besides, TGC of systems with $P = 6.0$ MPa was two times higher than that of systems with $P = 3.0$ MPa. In complement with the data of TGC, normalized gas consumption (NGC) was calculated with TGC and the moles of added TBAC (n_{TBAC}) which was collected with the test solutions, as shown in eqn (1). Its unit was $\text{molCH}_4 \text{ mol}^{-1} \text{TBAC}$.

$$\text{NGC} = \text{TGC}/n_{\text{TBAC}} \quad (1)$$

Different from the calculation of NGC with moles of consumed gas and water reported in refs.^{46,47} the calculation of NGC in the present study was proposed due to the fact that the amount of TBAC was the main factor affecting the hydrate forming amount in our systems. According to the structural parameters ($10\text{D}\cdot5\text{TBAC}\cdot150\text{H}_2\text{O}$, $V = 7.048 \times 10^{-27} \text{ m}^3$) reported by Rodionova *et al.*,³ the storage capacity of (TBAC + CH_4) hydrate should be $2 \text{ molCH}_4 \text{ mol}^{-1} \text{TBAC}$ (namely $52.768 \text{ m}^{-3} \text{CH}_4 \text{ m}^3 \text{SCH}$)⁴⁸ with TS-I structure. It should be noted that the NGC of A2 was higher than $2 \text{ molCH}_4 \text{ mol}^{-1} \text{TBAC}$. This might be caused by the structure of SCH, which would be checked and discussed in the next Section 3.2. From Fig. 5, it could be seen that the effects of P and T on NGC were similar to those on TGC. But considering the effect of w , it was greatly different. With the increase of w , NGC continuously decreased. Considering the fact that in all systems only double (TBAC + CH_4) SCH was formed, this phenomenon revealed that with the increase of w , the amount of CH_4 trapped in the unit cell of SCH continuously decreased, indicating that the cage occupancy decreased. Thus the system with high w , showing good stabilization effect in thermodynamics, did not perform well in gas storage, and the system with low salt concentration was a good choice for advanced gas storage. Besides, Fig. 5 illustrated that the gas consumption parameters were affected more by pressure than by temperature, thus for systems with same w , pressure was a key parameter for gas consumption. Since the gas consumption was affected by the experimental conditions, systems with TBAB reported in open literature could not be compared with the systems in this study. Thus the comparison of systems with different SCH formers under the same experimental conditions would be the subject of future study.

Furthermore, the rapid growth stage of hydrate growth process was studied in addition to understanding the hydrate formation kinetics. The process, shown as section C in Fig. 2, began from the end of the nucleation and ended at the point in which the slope of the plotted gas consumption- t curve sharply changed. In this section, the hydrate was formed evenly and quickly, represented by the constant slope of (gas consumption- t) curve. Thus the rapid growth time (RGT) and rapid growth rate (RGR) were chosen to illustrate this stage. As shown in Fig. 6, the RGT of systems with $P = 6.0$ MPa and $\Delta T = 6$ K were relatively long under different w , while in systems with $P = 3.0$ MPa, the RGT became shorter with the increase of w . Besides, the RGR was also an important parameter. As



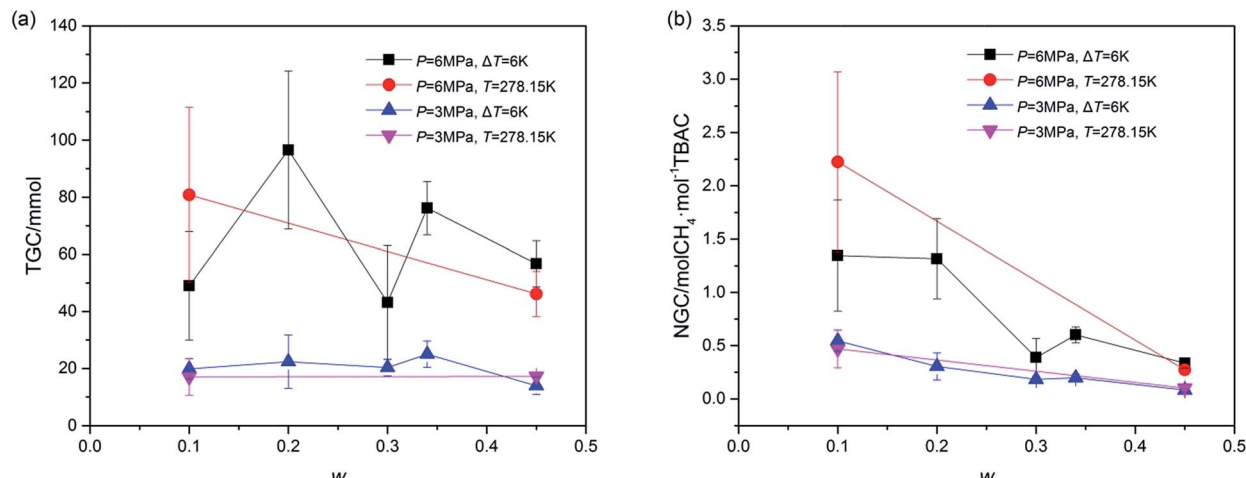


Fig. 5 Gas consumption (GC) of TBAC + CH₄ hydrate growth process at various w with different conditions: (a), total gas consumption (TGC); (b), normalized gas consumption (NGC). The error bars indicated the standard deviations of each system.

illustrated in Fig. 6, the RGR decreased vary slightly with the increase of w in each system. And the effect of T on RGR was larger in systems with $P = 6.0$ MPa than that in systems with $P = 3.0$ MPa. Fig. 6 revealed that the hydrate rapid growth process, different from gas consumption parameters, was affected more by temperature than by pressure. In other words, to improve the gas rapid growth process, temperature was a key parameter. In addition, these data suggested that the under the conditions of $\Delta T = 6$ K and $T = 278.15$ K, the running time for hydrate growth would be better within (40 to 100) min and (30 to 60) min, respectively.

3.2 Structural analysis of (TBAC + CH₄) hydrate

To understand the structure and molecular behavior of (TBAC + CH₄) SCHs, Raman measurement was carried out. Fig. 7 presented the Raman spectra of (TBAC + CH₄) hydrate samples. It could be seen that one peak appeared at 2912 cm⁻¹. In 2016, Jin

et al.⁴⁹ reported the Raman spectra of TBAB hydrate. The peak around 2912 cm⁻¹, corresponding to C-H vibration of TBA⁺, was much lower than the left peak and the right peak. While in this study, for systems with $w = (0.10$ and $0.20)$, the peak around 2912 cm⁻¹ was higher than other peaks. For systems with $w = (0.30$, 0.34 and $0.45)$, the peak at 2912 cm⁻¹ was higher than the left peak and almost as high as the right peak. Thus the peak at 2912 cm⁻¹ could be assigned to the C-H vibration of CH₄ molecules in 5¹² cages. This implied that small cages of TBAC SCHs were vacant to trap gas molecules and the cage occupancy decreased with the increase of w . It was well-known that CH₄ hydrate was formed based on structure I with two peaks at 2905 cm⁻¹ and 2915 cm⁻¹, corresponding to the CH₄ in large and small cages, respectively. The slight shift in wavenumbers (2915 cm⁻¹ to 2912 cm⁻¹) could be caused by the structure of double (TBAC + CH₄) SCH, in which the cation was engaged in the host lattice and affected the Raman spectra. Besides, it

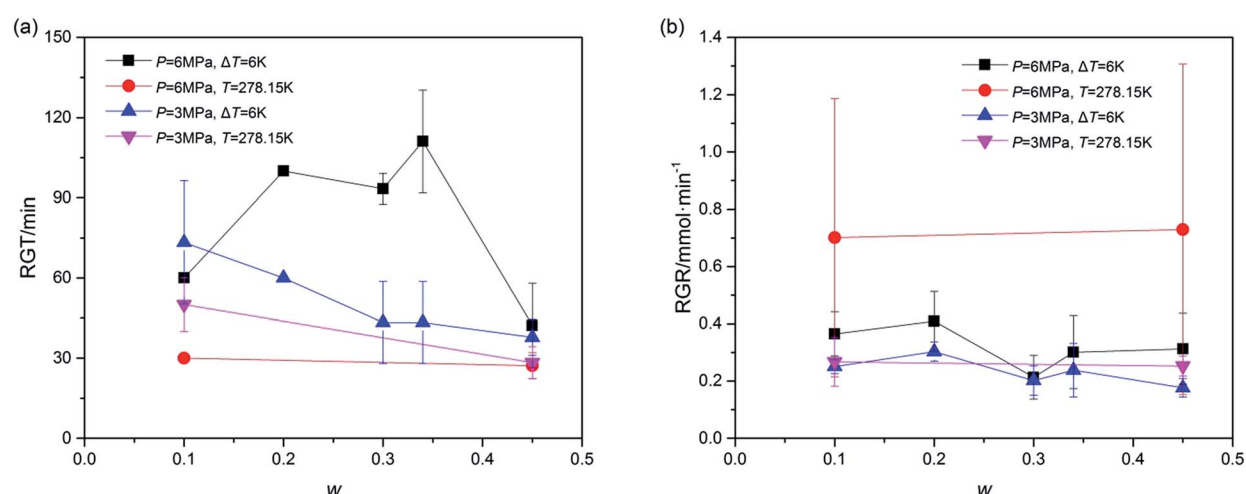


Fig. 6 Rapid growth process of TBAC + CH₄ hydrate: (a), rapid growth time (RGT); (b), rapid growth rate (RGR). The error bars indicated the standard deviations of each system.

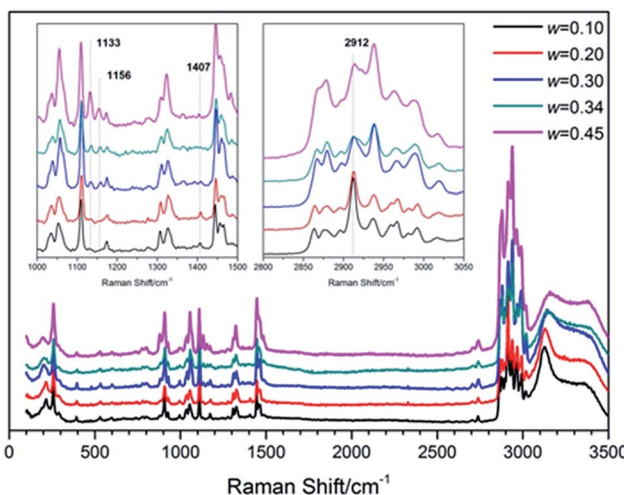


Fig. 7 Raman spectra of (TBAC + CH₄) SCHs formed with various w .

should be noted that this phenomenon was not the first time to show up. In 2015, Kim *et al.*^{39,40} studied the spectra of (TBAC + CO₂) SCHs and (TBAC + CO₂ + N₂) SCHs, and the results showed that the addition of TBAC caused a slight shift of wavenumbers from 1275 cm⁻¹ to 1272 cm⁻¹. Moreover, Hashimoto⁵⁰ provided the differences between HS-I SCHs and TS-I SCHs in Raman spectra in 2008. The differences mainly showed in the peaks around 1100 cm⁻¹ and 1400 cm⁻¹, and the latter one only showed up in HS-I. In our experimental systems, the peaks appeared at 1133 cm⁻¹, 1156 cm⁻¹ and 1407 cm⁻¹. And the peaks at 1133 cm⁻¹ and 1156 cm⁻¹ became more obvious with w increased. This result revealed that the (TBAC + CH₄) SCHs were formed on the base of HS-I with w = (0.10 and 0.20), TS-I with w = (0.30 and 0.34), and both (HS-I + TS-I) with w = 0.45, respectively. It was reported that pure TBAC hydrate was formed only on the basis of TS-I.³ Thus it was assumed that at low salt concentrations the addition of CH₄ induced the formation of HS-I since it had three D cages per TBAC which was higher than that of TS-I.

4. Conclusions

In this study, the formation kinetics of TBAC + CH₄ SCHs have been investigated with an isobaric method for potential application of gas storage at moderate temperatures. In order to fully understand the effect of different operation conditions, the experiment was carried out for systems with w = (0.10, 0.20, 0.30, 0.34 and 0.45) and P = (6.0 and 3.0) MPa at constant (ΔT or T). The IT became shorter with higher supersaturation environment, represented by higher P or lower T . The NGC continuously decreased with the increase of w , showing that the amount of CH₄ trapped in the 5¹² cages decreased rather than kept constant. The parameters of rapid growth stage, RGT and RGR, were studied to provide information for industrial operation. The suggested operation time for systems with ΔT = 6 K or T = 278.15 K was (40 to 100) min and (30 to 60) min, respectively. In addition, Raman spectra were employed to

investigated the structure of (TBAC + CH₄) SCHs. The spectra revealed that the SCHs were formed with hexagonal structure or tetragonal structure under different salt concentrations, which were different from the structures of pure TBAC hydrate. It was assumed that at low salt concentrations the addition of CH₄ induced the formation of hexagonal structure since it had three 5¹² cages per TBAC which was higher than that of tetragonal structure. These results were of great fundamental and practical importance for advanced hydrate-based gas storage.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors acknowledge the support from the National Natural Science Foundation of China (51606198 and 51376182), the Natural Science Foundation of Guangdong Province (2016A030310126), and Fund of Key Laboratory of Gas Hydrate Chinese Academy of Sciences (Y607jb1001).

References

- 1 E. D. Sloan and C. A. Koh, *Clathrate Hydrates of Natural Gases*, CRC Press (Taylor & Francis Group), 3rd edn, 2008.
- 2 W. Shimada, M. Shiro, H. Kondo, S. Takeya, H. Oyama, T. Ebinuma and H. Narita, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 2005, **61**, O65–O66.
- 3 T. V. Rodionova, V. Y. Komarov, G. V. Villevaeld, T. D. Karpova, N. V. Kuratieva and A. Y. Manakov, *J. Phys. Chem. B*, 2013, **117**, 10677–10685.
- 4 T. Rodionova, V. Komarov, G. Villevaeld, L. Aladko, T. Karpova and A. Manakov, *J. Phys. Chem. B*, 2010, **114**, 11838–11846.
- 5 V. Y. Komarov, T. V. Rodionova, I. S. Terekhova and N. V. Kuratieva, *J. Inclusion Phenom. Macrocyclic Chem.*, 2007, **59**, 11–15.
- 6 Y. Youn, J. Seol, M. Cha, Y.-H. Ahn and H. Lee, *J. Chem. Eng. Data*, 2014, **59**, 2004–2012.
- 7 J. W. Du and L. G. Wang, *Ind. Eng. Chem. Res.*, 2014, **53**, 1234–1241.
- 8 J. W. Du, H. J. Li and L. G. Wang, *Ind. Eng. Chem. Res.*, 2014, **53**, 8182–8187.
- 9 Z. X. Liao, X. Q. Guo, Y. Y. Zhao, Y. W. Wang, Q. Sun, A. X. Liu, C. Y. Sun and G. J. Chen, *Ind. Eng. Chem. Res.*, 2013, **52**, 18440–18446.
- 10 J. W. Du, D. Q. Liang, D. L. Li, Y. F. Chen and X. J. Li, *Ind. Eng. Chem. Res.*, 2011, **50**, 11720–11723.
- 11 X.-S. Li, Z.-M. Xia, Z.-Y. Chen, K.-F. Yan, G. Li and H.-J. Wu, *Ind. Eng. Chem. Res.*, 2010, **49**, 11614–11619.
- 12 Q. A. Sun, X. Q. Guo, A. X. Liu, B. Liu, Y. S. Huo and G. Y. Chen, *Ind. Eng. Chem. Res.*, 2011, **50**, 2284–2288.
- 13 P. Babu, W. I. Chin, R. Kumar and P. Linga, *Ind. Eng. Chem. Res.*, 2014, **53**, 4878–4887.
- 14 N. Ye, P. Zhang and Q. S. Liu, *Ind. Eng. Chem. Res.*, 2014, **53**, 10249–10255.



15 D.-l. Zhong, Y. Ye, C. Yang, Y. Bian and K. Ding, *Ind. Eng. Chem. Res.*, 2012, **51**, 14806–14813.

16 M. Garcia, R. Marriott and M. A. Clarke, *Ind. Eng. Chem. Res.*, 2016, **55**, 777–787.

17 S. Muromachi, R. Kamo, T. Abe, T. Hiaki and S. Takeya, *RSC Adv.*, 2017, **7**, 13590–13594.

18 H. Sakamoto, K. Sato, K. Shiraiwa, S. Takeya, M. Nakajima and R. Ohmura, *RSC Adv.*, 2011, **1**, 315–322.

19 C.-G. Xu and X.-S. Li, *RSC Adv.*, 2015, **5**, 54672–54699.

20 C.-G. Xu and X.-S. Li, *RSC Adv.*, 2014, **4**, 18301–18316.

21 P. T. Ngema, C. Petticrew, P. Naidoo, A. H. Mohammadi and D. Ramjugernath, *J. Chem. Eng. Data*, 2014, **59**, 466–475.

22 T. Tanii, *J. Jpn. Soc. Mech. Eng.*, 1994, **97**, 925–927.

23 X. Wang, M. Dennis and L. Hou, *Renewable Sustainable Energy Rev.*, 2014, **36**, 34–51.

24 S. Babaee, H. Hashemi, A. H. Mohammadi, P. Naidoo and D. Ramjugernath, *J. Chem. Thermodyn.*, 2015, **81**, 52–59.

25 G. Li, D. Liu and Y. Xie, *J. Therm. Anal. Calorim.*, 2010, **102**, 819–826.

26 D. L. Fowler, W. V. Loebenstein, D. B. Pall and C. A. Kraus, *J. Am. Chem. Soc.*, 1940, **62**, 1140–1142.

27 G. A. Jeffrey, *Acc. Chem. Res.*, 1969, **2**, 344–352.

28 D. Mech, G. Pandey and J. S. Sangwai, *Fluid Phase Equilibr.*, 2015, **402**, 9–17.

29 D. Mech and J. S. Sangwai, *J. Chem. Eng. Data*, 2014, **59**, 3932–3937.

30 D. Mech and J. S. Sangwai, *J. Chem. Eng. Data*, 2016, **61**, 3607–3617.

31 J. S. Sangwai and L. Oellrich, *Fluid Phase Equilibr.*, 2014, **367**, 95–102.

32 H. Najibi, K. Momeni, M. T. Sadeghi and A. H. Mohammadi, *J. Chem. Thermodyn.*, 2015, **87**, 122–128.

33 Q. Sun, J. Liu, A. Liu, X. Guo, L. Yang and J. Zhang, *Int. J. Hydrogen Energy*, 2015, **40**, 6358–6364.

34 S. Muromachi, K. A. Udachin, S. Alavi, R. Ohmura and J. A. Ripmeester, *Chem. Commun.*, 2016, **52**, 5621–5624.

35 W. Lee, Y.-S. Kim and S.-P. Kang, *Chem. Eng. J.*, 2018, **331**, 1–7.

36 H. Roosta, S. Khosharay and F. Varaminian, *Energy Convers. Manage.*, 2013, **76**, 499–505.

37 J. Verrett and P. Servio, *Can. J. Chem. Eng.*, 2016, **94**, 2138–2144.

38 L.-L. Shi and D.-Q. Liang, *J. Chem. Eng. Data*, 2014, **59**, 3705–3709.

39 S. Kim, I.-H. Baek, J.-K. You and Y. Seo, *Appl. Energy*, 2015, **140**, 107–112.

40 S. Kim and Y. Seo, *Appl. Energy*, 2015, **154**, 987–994.

41 M. Oshima, M. Kida and J. Nagao, *J. Chem. Eng. Data*, 2016, **61**, 3334–3340.

42 L. Shi, L. Yi, X. Shen, W. Wu and D. Liang, *J. Mol. Liq.*, 2017, **229**, 98–105.

43 T. Makino, T. Yamamoto, K. Nagata, H. Sakamoto, S. Hashimoto, T. Sugahara and K. Ohgaki, *J. Chem. Eng. Data*, 2010, **55**, 839–841.

44 Z.-G. Sun and C.-G. Liu, *J. Chem. Eng. Data*, 2012, **57**, 978–981.

45 L.-l. Shi and D.-q. Liang, *Fluid Phase Equilibr.*, 2015, **386**, 149–154.

46 H. P. Veluswamy, W. J. Ang, D. Zhao and P. Linga, *Chem. Eng. Sci.*, 2015, **132**, 186–199.

47 H. P. Veluswamy, A. Kumar, R. Kumar and P. Linga, *Appl. Energy*, 2017, **188**, 190–199.

48 D. Mech, P. Gupta and J. S. Sangwai, *J. Nat. Gas Sci. Eng.*, 2016, **35**, 1519–1534.

49 Y. Jin, M. Kida and J. Nagao, *J. Chem. Eng. Data*, 2016, **61**, 679–685.

50 S. Hashimoto, T. Sugahara, M. Moritoki, H. Sato and K. Ohgaki, *Chem. Eng. Sci.*, 2008, **63**, 1092–1097.

