The Relationship between Thermal Characteristics and Microstructure/Composition of Carbon Dioxide Hydrate in the Presence of Cyclopentane

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Abstract: Hydrate-based carbon dioxide (CO$_2$) separation and capture is a new technology for achieving CO$_2$ emission reduction. However, it is still not commercially applied for the ambiguity of microscopic hydrate formation mechanism. In a constant volume experiment of hydrate formation, there are two or more pressure platforms, indicating that there might be two or more different hydrates formation in succession. In order to reveal the relationship between the microscopic process and the gas consumption in the process of hydrate formation, hydrate composition and formation mechanism of cyclopentane-CO$_2$ (CP-CO$_2$) system was investigated using a differential scanning calorimeter (DSC) and Raman spectroscopy. The results indicated CO$_2$-CP binary hydrate and CO$_2$ hydrate are formed successively, and they coexist in the final hydrate. CP-CO$_2$ binary hydrates forms preferentially, and as crystal seeds, inducing the formation of CO$_2$ hydrates. The two hydrates formation processes cause the two pressure-drops. The results provide a scientific basis for increasing the gas consumption in different stages of gas hydrate formation in the presence of hydrate formation promoter.

Keywords: carbon emission reduction; hydrate; microscopic mechanism; binary hydrate; DSC

1. Introduction

Energy and environment issues are the focus of researchers all over the world. With the continuous development of industry, the CO$_2$ emissions from fossil fuel combustion increase year by year, which has become the main reason for greenhouse effect such as global warming [1]. As there is no new energy that can completely replace the fossil energy to social development, fossil energy will continue to be the main energy to support economic development, thus the sustainable utilization of fossil energy is of great significance [2]. Therefore, while maintaining the efficient and sustainable utilization of fossil energy, it is very important to separate and capture CO$_2$ from the exhaust gas [3]. Many meaningful methods of atmospheric CO$_2$ fixation in inorganic and biomimetic structures have been proposed [4]. The traditional methods of CO$_2$ separation include physical absorption, chemical adsorption, membrane separation, cryogenic separation and chemical cycle combustion method, but all these methods have their individual defects, for instance, most
of the chemical adsorbents aren’t environmental friendly, the capacity of a membrane is small and the membrane is consumable, therefore, it is expected to develop new and environmental friendly and efficient technology for gas separation [5–7]. Hydrate-based gas separation technology has attracted great attention because of its advantages of simple process, low energy consumption and relatively environmental friendly [8,9].

Gas hydrate is a kind of non-stoichiometric crystalline compound, which is composed of water molecules as the host molecules and gas molecules (CH₄, CO₂, H₂, etc.) as the guest molecules. Water molecules form cages through hydrogen bond, and gas molecules enter the cages and make the cages more stable by Van der Waals force [10]. Depending on the pressure and the molecular properties of the guest, gas hydrates usually have different structures. The three main structures include cubic structure I (sI), cubic structure II (sII) and hexagonal structure H (sH), and sI hydrate crystal unit is composed of two 5₁² and six 5₁²6₂ cages with chemical formula of 2 [5₁²] · 6 [5₁²6₂] · 46H₂O. sII hydrate crystal unit is composed of sixteen 5₁² and eight 5₁²6₄ cages with chemical formula of 16 [5₁²] · 8 [5₁²6₄] · 136H₂O. sH hydrate crystal unit is composed of three 5₁² cages, two 4½5½₆₃ cages and one 5½₆₆ cages with chemical formula of 3 [5₁²] · 2 [4½5½₆₃] · 1 [5½₆₆] · 34H₂O [11,12]. The principle of hydrate-based gas separation from gas mixture is that the different gas components have their individual gas hydrate formation equilibrium conditions, and the component with relatively moderate equilibrium condition preferentially enters the hydrate cage and enriches in the hydrate phase, while the remaining components enrich in the residual gas phase. For example, compared with H₂ and N₂, the phase equilibrium pressure of CO₂ hydrate is much lower at a certain temperature, and it is easier to enter the hydrate to enrich in the hydrate, so as to achieve the purpose of separating CO₂ from CO₂/H₂ or CO₂/N₂ mixture [13]. In fact, gas hydrate formation could only happen under the condition of high pressure or and low temperature. For example, the phase equilibrium pressure is about 2.4 MPa for pure CO₂ forming CO₂ hydrate at 5 °C in pure water system. And the higher pressure or lower temperature means the higher energy consumption. Therefore, various promoters have been developed to moderate the hydrate formation conditions. All the promoters are divided into thermodynamic promoter and kinetic promoter [14]. The thermodynamic promoter itself, such as Tetrahydrofuran (THF) [15–18], CP [19], Tetrabutyl Ammonia Salt (TBA+) [20–23], Propane (C₃H₈) [24,25], can form hydrate with a relatively moderate phase equilibrium condition. Therefore, the presence of the thermodynamic promoter can moderate the hydrate formation temperature and pressure. Taken CP as an example, relative to CO₂ or CH₄, CP could form CP hydrate at quite higher temperature and lower pressure, thus on one hand, the CP hydrate which formed in advanced could offer cages for CO₂ or CH₄ to form CO₂-CP or CH₄-CP binary hydrate, on the other hand, the CP hydrate could induce the formation of CO₂ hydrate or CH₄ hydrate formation as crystal seeds, moderating the conditions of forming CO₂ hydrate or CH₄ hydrate [26,27]. The function of kinetic promoter, such as Sodium Dodecyl Sulfonate (SDS) [28,29], Tween 80 [30], Dimethyl Sulphoxide (DMSO) [31], is to improve the fugacity of gas in liquid phase by changing the surface tension and promoting gas dissolution. Currently, the main thermodynamic promoters include THF, CP and TBAB, etc. Many works have been carried out to investigate the effect of the promoters on hydrate formation equilibrium conditions, kinetics and thermodynamics of hydrate formation, gas separation efficiency, micro-mechanism of hydrate formation, etc. Without any questions, the positive effects of the thermodynamic promoters on hydrate formation have been systematically proven, and some new synergistic promoters such as Tetrahydrofuran bromide + Dodecyl Trimethyl Ammonium Chloride (TBAB+DTAC) [7], CP+TBAB [32], THF+SDS [28], TBAB+DMSO [31,33] have also been developed. However, as reported previously, different thermodynamic promoters have different selectivity for gases in the process of hydrate formation depending on their physical and chemical properties. For example, CP and THF are relatively more conducive to the formation of CO₂ and CH₄ hydrates, while TBAB could significantly reduce the formation conditions of hydrate but being disadvantage of the gas consumption for the hydrate agglomeration [14]. Even so,
the scientific basis supporting these experimental results, especially the thermodynamic micro-mechanism, is still unclear.

The thermodynamic micro-mechanism of gas hydrate formation is usually inferred by molecular dynamic (MD) simulation [34–39]. It could also be obtained by measuring the reaction heat of hydrate formation. Since the 1980s, differential scanning calorimetry (DSC) has been used to study the thermal properties of hydrates, such as hydrate decomposition enthalpy and heat capacity [40,41]. Under a certain temperature, it is a kind of analytical method for determining the energy difference related to the temperature between the delivered substance and the reference substance. The gas hydrate decomposition enthalpy ($\Delta H_d$) is a key parameter for predicting the heat for hydrate decomposing into water and gas. Koh et al. [42] used DSC to measure the decomposition heat of hydrate formation by carrying out experiments on simulated natural gas hydrate (NGH) formation and decomposition under environmental pressure, and used DSC to quantify and compare the effects of various kinetic inhibitors. Dalmazzone et al. [43] employed high-pressure DSC to characterize the limits of the thermodynamic stability of methane hydrate and other natural gas hydrates in the presence of hydrate formation kinetic inhibitors (KIs). Zhang et al. [44] measured the decomposition temperature of $\text{H}_2 + \text{CP}$ and $\text{CO}_2 + \text{CP}$ in different pressures by high pressure DSC. The results showed that the decomposition temperature of $\text{H}_2$ hydrate depends on the properties of additives, while the decomposition temperature of $\text{CO}_2$ hydrate is independent of the properties of the additives. In addition, Seo et al. [45,46] (from Ulsan National Institute of Science and Technology) used HP $\mu$-DSC to prove the existence of sI and sII hydrates in the process of CH$_4$–CO$_2$ replacement and obtain the transformation enthalpy of hydrate structure from sH to sI in the CO$_2$ capture and sequestration by hydrate method, which are of great significance for further study on the mechanism of gas hydrate technology.

The hydrate formation or/and decomposition enthalpy is closely related to the hydrate structure and composition. And the hydrate structure and composition could be determined by many micro analytical methods, e.g., Raman Spectroscopy, Fourier Transform Infrared Spectroscopy (FTIR), Powder X-ray Diffraction (PXRD), Nuclear Magnetic Resonance (NMR) [47–49]. For instance, Chen et al. [50] investigated structures of CO$_2$ hydrate and THF + CO$_2$ binary hydrate by Raman spectra, and they found in the pure CO$_2$ hydrate, CO$_2$ molecules occupy both medium $5^{12}6^2$ and small $5^{12}$ cages, while CO$_2$ molecules only occupy the small $5^{12}$ cages in the binary hydrate. They further believed that THF occupies the medium $5^{12}6^4$ cages is the reason why the gas consumption in THF system is lower than that in pure water system. Xu et al. [14] successfully achieved the mechanism of hydrate formation promoters affecting on the gas selectivity in the process of hydrate formation by Raman and Cryo-SEM. Seo et al. [51] carried out the experiments to form CH$_4$ hydrate in the presence of tetra-iso-amyl ammonium bromide (TiAAB), and investigated the relationship between the transformation of hydrate structure among sI, sII or/and sH and the composition of gas hydrate by DSC and Raman, then they found the inclusion of CH$_4$ had few effects on the semi-clathrate structure.

Up to now, there is still no authoritative consensus on the micro process of gas hydrate formation, especially whether there is a correlation between the thermodynamic properties of hydrate formation and the hydrate structure and composition. In order to further reveal the relationship between them, the thermodynamic properties of CO$_2$ hydrate formation in the presence of CP were studied by DSC, and the hydrate composition and structure were analyzed by Raman. Through the scientific comparison and analysis of the experimental results, it is expected to find the corresponding relationship between the formation thermodynamic properties of gas hydrate and the hydrate structure and composition.

2. Experimental
2.1. Experimental Apparatus and Materials

The low-temperature and high-pressure DSC and Raman spectrometer used in this experiment are Setaram BT 2.15 and Horiba LabRam, respectively. CO$_2$ (A.R. > 99.99%) is pro-
vided by Guangzhou Shengying Gas Co., Ltd. (Guangzhou, China), and CP (A.R. > 99.0%) is offered by Xiamen Pioneer Chemical Reagent Co., Ltd. (Xiamen, China), and the deionized water with a resistivity of 18.25 mΩ/cm is self-produced using an ultra-pure water system supplied by Nanjing Ultrapure Water Technology Co., Ltd., Nanjing, China.

The whole experimental schematic diagram is shown in Figure 1. As shown, there are two sets of experimental equipment. The left is special for gas hydrate formation and Raman measurement, and the right is special for DSC measurement. For the left, the high-pressure cylinder reactor is made of 304 stainless steel with an internal volume of 100 mL and a maximum bearing pressure of 10 MPa. There are visual windows at the front and back of the reactor, through which the hydrate formation and decomposition process can be observed. There is a flange on the reactor and it is connected with the gas cylinder through the conduit. The temperature and pressure in the reactor are measured by a Omega Pt 100 thermocouple with the uncertainties of 0.1 °C and a Trafag NAH 8253 pressure transmitter with accuracy of 0.3%. Temperature and pressure are directly transferred to computer through data acquisition system. The reactor is placed on a magnetic stirrer with a rotating speed of 0–2000 r/min. The material in the reactor can be fully mixed by the magnetic stirrer. The system temperature is regulated by a water-bath jacket outside the reactor, and the jacket is filled with a 25 wt% glycol aqueous solution. DSC consists of three main parts: gas control platform, reaction tank and calorimetric system, liquid nitrogen (LN) temperature control system. An air-bath tank is set out around the DSC, which is full of N₂ during the measurement, and a LN tank is set in the DSC, which is filled with LN. A DSC cell with inner volume of 3.60 mL is placed in the LN tank for test. Here is a program controller for setting the temperature and pressure program.

![Schematic diagram of the experimental equipment.](image)

**Figure 1.** Schematic diagram of the experimental equipment.

### 2.2. Experimental Procedure

#### 2.2.1. DSC Experiment

The process of the DSC experiment is similar to that reported in Reference [32]. Before the experiment, the cell is firstly rinsed with the ionized water at least 3 times, and then thoroughly dried with a dryer. After being injected about 1.0 g solution (CP solution 5 v%), the cell is transferred to the He-bath chamber. The chamber is vacuumized to −1.0 MPa, and then is pumped to 1.0 MPa with pure helium (He). The feed gas is injected into the cell to exclude the air and then pumped to the set experimental pressure with an automatic pressurizer. Whereafter, the temperature program is set according to the experimental scheme, and then started. For each experiment, the same program runs four times automatically. The detail procedure of DSC experiments is set as shown in Table 1.
In this experiment, the final sample is 1.0550 g. And then install the high-pressure sample cell into the measuring unit according to the operation specification.

| Exp. Run | P (MPa) | T (°C)       | Cooling Rate (K·min⁻¹) | Constant Time (h) | Heating Rate (K·min⁻¹) |
|----------|---------|--------------|------------------------|------------------|------------------------|
| 1        | 2.5     | 20.0~−5.0    | 0.02                   | 5                | 0.02                   |
| 2        | 2.5     | 20.0~−10.0   | 0.02                   | 5                | 0.02                   |
| 3        | 2.5     | 20.0~−8.8    | 0.02                   | 5                | 0.02                   |
| 4        | 5.0     | 20.0~−8.8    | 0.02                   | 5                | 0.02                   |
| 5        | 0       | 20.0~−8.8    | 0.02                   | 5                | 0.02                   |
| 6        | 3.5     | 20.0~−8.8    | 0.02                   | 5                | 0.02                   |
| 7        | 6.0     | 20.0~−8.8    | 0.02                   | 5                | 0.02                   |

2.2.2. Raman Experiment

The steps of Raman experiment are similar to those in reference [52], including hydrate formation, hydrate sampling, Raman measurement, etc., but due to the update of the test system, the Raman spectrometer was replaced by InVia Reflex from Renishaw with a single monochromator of 1800 grooves/mm grating, and a multichannel air-cooled charge coupled device (CCD) detector is employed to measure the simulated CH₄-CO₂ binary hydrate, using a 532 nm Argon laser with 100 mW. Before the measurement, the Raman spectrum is calibrated with standard silicon (Si) crystal of 520.7 cm⁻¹.

All the DSC measurement and Raman measurement were repeated by 3 times in order to ensure all measurements were repeatable. In addition, the pressure in the hydrate formation experiment and Raman measurement were gauge pressure, but the pressure in the DSC measurement was absolute pressure.

3. Results and Discussion

3.1. Analysis of Hydrate Formation Process

First of all, two groups of gas hydrate formation experiments were carried out under different temperature and pressure conditions, and in each group, the same experiment was repeated 3 times. Figure 2 shows the changes of temperature and pressure in the process of CO₂ hydrate formation in the presence of CP with initial pressure of 2.5 MPa. Same as other reported figures of pressure-temperature vs. time in the hydrate formation processes, the pressure drops sharply firstly and then becomes stable gradually, and the temperature also remains nearly the same. However, there is a little temperature rise in the time zone where the pressure drops sharply. Actually, the sharp pressure-drop could be attributed to two stages. One is the dissolution of gas, followed by the gas hydrates formation. Because the solubility of gas in solution is very fixed under certain temperature and pressure, especially for those gases with poor solubility or low solubility, the solubility of gas is not the main contributor to pressure drop compared with gas entering hydrate cage. Therefore, it is feasible to use the real gas equation to calculate the gas consumption in the process of gas hydrate formation and then calculate the gas consumption rate to further characterize the relevant hydrate formation kinetics. The equation is shown as following: [52]

\[
\Delta n = \left( \frac{PV}{zRT} \right)_f - \left( \frac{PV}{zRT} \right)_0
\]

where “\(\Delta n\)” is the mole of gas consumed in the hydrate formation or entering the hydrate slurry, and “\(P\)”, “\(V\)”, “\(T\)” are the pressure, volume, temperature of the gas phase in the reactor, respectively. In this work, it is assumed the liquid and hydrate are incompressible. “\(R\)” is the gas constant, and “\(z\)” is the compressibility factor. The subscript “\(0\)” and “\(f\)” indicate the starting time and ending time of the hydrate formation, respectively.
As shown in Figure 2, the total gas consumption per mole water is 0.0187 mol$^{-1}$, and there are two obvious pressure platforms (I and II) over time. According to the trend of the pressure-drop in the figure, the whole pressure drop process is divided into two stages, the first stage (stage I) and the second stage (stage II) with two platforms as the boundary, respectively. The stage I, from 0 to 174 min, is defined as the first hydrate formation platform. And at this stage, there are two processes, including gas dissolution followed by the hydrate formation. The second stage, from the 174th minute to the end of hydrate formation, is the hydrate secondary formation platform. At this stage, a large amount of gas is consumed, but the average gas consumption is relatively slower than that at the stage I. The second platform indicates the existence of hydrate secondary generation in the system. Figure 3 shows the temperature and pressure changes over time in the process of CO$_2$ hydrate formation in the presence of CP with initial pressure of 5.0 MPa. As shown, the general trend of pressure change is similar to Figure 2, and there are two pressure platforms and the pressure-drop rate at the stage II is slower than that at the stage I. Therefore, it also shows that there is a secondary hydrate formation at 5.0 MPa. Unlike Figure 2, the first stage in Figure 3 has a higher pressure drop rate and a shorter time for the second pressure to fall to steady. Additionally different from Figure 2, the pressure drop rate of the first stage in Figure 3 is higher, and the time for the second pressure to fall to stability is shorter, which shows that the increase of pressure is more conducive to the first formation of gas hydrate. In fact, in a fixed system, the pressure-drop is completely caused by the gas consumption in the process of gas hydrate formation. Therefore, in the later content, we use gas consumption instead of pressure-drop. As our previous study [53], the gas consumption at the second stage is slower than that at the first stage, which is mainly due to the hydrate formed at the first stage hinders the gas diffusion at the second stage. However, the diagram only reflects the P-V-T change rule of hydrate formation process, so it can only judge the existence of secondary formation in the whole process, but it is not clear about the information of hydrate structure, composition and thermodynamic characteristics in the hydrate secondary formation. We combine DSC experiment with Raman analysis to obtain these information and show it in the following content.
enthalpy and decomposition enthalpy of the same gas hydrate is 300–400. The heat of formation and melting enthalpy are very close. The ratio of hydrate total decomposition enthalpy to hydrate enthalpy is 99.26%, 101.52%, 99.48%, 100.23%, 101.46% and 97.15%, respectively, indicating that hydrate formation and decomposition in the system are in accordance with heat conservation. For each experiment as run 2–4, there are two distinct formation peaks and decomposition peaks, and in the time dimension, the earlier formation peak corresponds to the later decomposition peak, while the later formation peak corresponds to the earlier decomposition peak. The absolute value of the enthalpy corresponding to the formation or decomposition peaks is almost the same, but the enthalpy values of the two decomposition peaks are quite different, which indicates that two different types of hydrates have been formed in the process. This result is consistent with the results shown in Figures 2 and 3.

Figure 3. The CP-CO$_2$ system pressure and temperature curves at 5.0 MPa.

3.2. DSC and Raman Measurement

Figure 4 shows the heat flow and temperature changes vs. time corresponding to the ice formation and melting under atmosphere with the Setaram BT 2.15. As shown, the ice starts to form at about $-11.20\, ^\circ$C and melt at 0.05 $^\circ$C and the corresponding formation enthalpy and melting enthalpy are $-335.29$ and $334.07$ J/g. Because enthalpy is an integral value obtained by the software program through calculating the area of the curve integral, there will inevitably be errors in the integration process, and as long as the error is less than 5%, we think the integral value is reasonable. Generally, in the system, the absolute value of the formation enthalpy and decomposition enthalpy of the same gas hydrate is equal, even if the integration error is considered, the error should be less than 5%. Figures 5–11 show the heat flow and temperature changes vs. time corresponding to the experiments listed in Table 1, and Table 2 shows the heat flow and temperature of the peaks corresponding to hydrate formation and dissociation under different experimental runs. In each figure, the black line shows that the sample temperature changes with time with the DSC temperature program, while the red line represents the heat flow changes with time. As shown, there are two exothermic peaks corresponding to hydrate dissociation in each figure, which means there do be two kinds of compounds formed in the system. Moreover, for each figure, the absolute values of hydrate formation enthalpy and decomposition enthalpy are very close. The ratio of hydrate total decomposition enthalpy to hydrate formation enthalpy is 99.26%, 101.52%, 99.48%, 100.23%, 101.46% and 97.15%, respectively, indicating that hydrate formation and decomposition in the system are in accordance with heat conservation. For each experiment as run 2–4, there are two distinct formation peaks and decomposition peaks, and in the time dimension, the earlier formation peak corresponds to the later decomposition peak, while the later formation peak corresponds to the earlier decomposition peak. The absolute value of the enthalpy corresponding to the formation or decomposition peaks is almost the same, but the enthalpy values of the two decomposition peaks are quite different, which indicates that two different types of hydrates have been formed in the process. This result is consistent with the results shown in Figures 2 and 3.
Table 2. The heat flow and temperature of the peaks corresponding to hydrate formation and dissociation under different experimental runs.

| Exp. Run | P (MPa) | T (°C) | First Formation Peak | Second Formation Peak | First Decomposition Peak | Second Decomposition Peak | ∆Hf,1/∆Hf,2 (%) |
|----------|---------|--------|----------------------|-----------------------|-------------------------|--------------------------|------------------|
|          |         |        | T (°C)    | ΔHf (J·g⁻¹) | T (°C)    | ΔHf (J·g⁻¹) | T (°C)    | ΔHf (J·g⁻¹) | T (°C)    | ΔHf (J·g⁻¹) |
| 1        | 2.5     | 20.0   | −5.0      | 0.34 −50.823 | −      | −          | 5.80      | 37.211     | 6.93      | 13.234     | 99.26 |
| 2        | 2.5     | 20.0   | −10.0     | 3.66 −63.97 | −6.19 | −217.557  | −1.72 | 214.034   | 5.71      | 67.212     | 99.48 |
| 3        | 2.5     | 20.0   | −8.8      | 3.65 −67.769 | −6.50 | 243.679  | −1.72 | 242.622   | 5.70      | 67.212     | 99.48 |
| 4        | 5.0     | 20.0   | −8.8      | 13.98 −77.013 | 3.21 | −39.510  | 10.03     | 39.014    | 14.36     | 77.775     | 100.23 |
| 5        | 0       | 20.0   | −8.8      | −2.63 −0.533 | 1.54 | 219.444  | 4.85      | 118.254   | 101.46    | 101.46    | 101.46 |
| 6        | 3.5     | 20.0   | −8.8      | 0.607 −113.919 | −5.90 | −218.931 | 1.54      | 219.444   | 4.85      | 118.254   | 101.46 |
| 7        | 6.0     | 20.0   | −8.8      | T: 6.97     | −162 | 17.502   | 10.216    | 438.734   | 97.15     | 97.15     |

Remark: P, T, ΔHf, ΔHd, ΔHf,1, ΔHf,2 presents operation pressure, sample temperature in the DSC cell, hydrate formation enthalpy, hydrate decomposition enthalpy, total hydrate formation enthalpy and total hydrate decomposition enthalpy, respectively.

Figure 4. The heat flow and temperature changes vs. time corresponding to the ice formation and melting with the Setaram BT 2.15.

During the hydrate formation in the static water system, the decomposition enthalpy may vary greatly under different experimental conditions because of the effects of factors such as interference of ice, the wall-climbing of hydrate and the difficulty of gas diffusion in hydrate. Even so, the gas molecular occupancy in hydrate cage can be qualitatively analyzed according to the absolute value of enthalpy as long as the structure of hydrate is determined. The higher the hydrate decomposition enthalpy, the higher the gas occupancy in the hydrate cages. Temperature is an important index that can qualitatively determine the type of hydrate. In other words, it could be determined what kind of the hydrate is according to the hydrate decomposition temperature. As shown in Table 2 and Figure 5, the two decomposition peaks occur at around 5.80 and 6.93 °C, respectively. Both the temperatures are higher than freezing temperature, indicating that no ice forms in the hydrate is according to the hydrate decomposition temperature. As shown in Table 2 and Figure 5, the two decomposition peaks occur at around 5.80 and 6.93 °C, respectively. Both the temperatures are higher than freezing temperature, indicating that no ice forms in the experiment of run 1. Compared with the results of other experiments, the temperatures corresponding to the two decomposition peaks are relatively close. This is also reflected in the corresponding formation peaks, as shown in Figure 5. In fact, two hydrates have been formed successively, but only one obvious formation peak can be seen. The reason is that the formation temperatures of the two hydrates are almost the same, so that the corresponding peaks are overlapped and hard to be separated. According to the decomposition temperature, actually, the two temperatures are close, thus there might be two possibilities, one, there might be two different kinds of hydrates, CP-CO2 binary hydrate and CO2 hydrate, another, there might be two endothermic processes during the hydrate
decomposition that the two kinds of cages ($5^{12}$ and $5^{12}6^4$) in the hydrate break down in order.

![Figure 5](image1.png)

**Figure 5.** Heat flow and temperature curve at 20.0–5.0 °C and 5.0 MPa.

![Figure 6](image2.png)

**Figure 6.** Heat flow and temperature curve at 20.0–10.0 °C and 2.5 MPa.

Admittedly, this result is different from other experimental results. As shown in Table 1, The experimental programs of run 2 and 3 are same except for the lower-limit of temperature, and the results of the run 2 and 3 listed in Table 2 are almost same, indicating that the lower-limit of temperature of 8.8 °C is reasonable for the experiment and the lower temperature is no need. For the experiments of runs 2 and 3, as shown in Table 2 and Figures 6 and 7, on one hand, the temperatures of the two decomposition peaks are around −1.72 and 5.71 °C, respectively, which are different very much. The first decomposition temperature is lower than the freezing point, while the second one is higher than the freezing point. Thus, it is certain that the second peak does not correspond to the ice melting. However, it is difficult to judge whether the first peak is ice peak by DSC result directly. For this reason, we have measured the ice point of CP-water system under atmospheric pressure. The temperature procedure of the DSC measurement is shown in Table 1 as Exp. Run 5, and the results of the DSC measurement are shown in Table 2.
and Figure 9. As shown, there are a very sharp exothermic peak at $-1.63 \, ^\circ C$, a wide and irregular endothermic peak at $-1.50 \, ^\circ C$, and two small exothermic peaks before and after the endothermic peak. Compared with other DSC results, it can be seen that the heat flow values corresponding to the exothermic and endothermic peaks in Figure 9 are quite small, that’s also the reason there are two small exothermic peaks before and after the endothermic peak. The absence of the secondary formation in other figures does not mean that there is no hydrate secondary formation, but that the heat flow value of hydrate decomposition is larger, so that the heat flow of the smaller hydrate secondary formation is negligible in the figure. In addition, we carried out an experiment to measure the freezing point of CP solution at atmospheric pressure. In the experiment, the reactor was injected with only 30 mL CP emulsion (5 v%) but no gas, then the system cooled gradually. As shown in Figure 12, the CP emulsion began to transfer into solid phase at around $-1.50 \, ^\circ C$. And about 2 h later, almost all CP emulsions were converted to solid. The solid is a mixture of CP hydrate and ice. According to the law of thermodynamics, the higher the pressure, the lower the freezing point of the liquid. Therefore, for Figures 6 and 7, based on the DSC results of this experiment, we conclude that the first peak presents the ice melting in the mixed system because the temperature is quite close to the freezing point, and the second peak presents the decomposition of CP-CO$_2$ binary hydrate because the temperature is quite close to that showed in Figure 5.

![Figure 7. Heat flow and temperature curve at 20.0 $\sim$ $-8.8 \, ^\circ C$ and 2.5 MPa.](image-url)

In order to further investigate the effect of the operation pressure on the thermodynamics of the hydrate formation, the experiment of run 4 was carried out under the constant pressure of 5.0 MPa. It is worth noting that in the DSC measurement process, the pressure of 5.0 MPa rather than higher pressure is selected to avoid CO$_2$ liquefaction. As shown in Table 2 and Figure 8, the two decomposition peaks with decomposition enthalpies of 39.01 and 77.78 J·g$^{-1}$ occur at 10.03 and 14.16 $^\circ C$, respectively. It can be determined that these two peaks are independent of ice due to their corresponding temperatures much higher than $-1.50 \, ^\circ C$. And the results are also quite different from those achieved in the experiments of run 1~3. However, from the perspective of decomposition enthalpy, the decomposition heat ($\Delta H_d$) corresponding to the first decomposition peak in Figure 8 is close to that in Figure 5, while the decomposition heat corresponding to the second decomposition peak is close to that in Figure 6. Therefore, it could be concluded that, firstly, the increase of pressure from 2.5 MPa to 5.0 MPa makes the decomposition temperature of hydrate increase significantly, secondly, it can be clearly determined that there are two different hydrates, and according to the composition of the substances in the system, it can be deduced that the first peak corresponds to CP-CO$_2$ binary hydrate, and the second de-
composition peak corresponds to CO$_2$ hydrate. The result is consistent with that obtained by Seo et al. [19]. In order to further clarify the influence of the pressure on the hydrate formation, we carried out another two runs of experiments (Exp. Run 6 and 7 in Table 1) with constant pressures of 3.5 MPa and 6.0 MPa, and the specific heat flow changes during hydrate formation and decomposition are shown in Figures 10 and 11. As shown, there are two decomposition peaks at $-1.54 \degree C$ and $4.85 \degree C$ in Figure 10 and $-1.62 \degree C$ and $10.22 \degree C$ in Figure 11, respectively. For the heat flow peak at around 4.85-6.00 \degree C under 3.5 MPa or less, what we can determine is the hydrate is CP-CO$_2$ binary hydrate because the Raman spectrum shows both CO$_2$ and CP are included in the hydrate, as shown in Figure 13. In Figure 13, it can be found two peaks appear at 1275.4 and 1379.5 cm$^{-1}$, and another two peaks appear at 2981.5 and 2875.0 cm$^{-1}$. The first two peaks are the characteristic peaks of CO$_2$, and the second two peaks correspond to CP. Thus, it could be inferred that CO$_2$ and CP coexist in the hydrate. However, for Figure 11, due to that the heat flow peak corresponding to hydrate decomposition at 10.22 \degree C is a relatively broad peak, its composition needs to be further determined by peak-differentiating and fitting according to the Gaussian-Lorentzian function [54].

![Figure 8. Heat flow and temperature curve at 20.0-8.8 \degree C and 5.0 MPa.](image)

As mentioned above, on one hand, compared the experimental results of runs 3 with run 6, the pressure rises from 2.5 MPa to 3.5 MPa did not affect the formation of ice and hydrate structure in the system, but only increased the temperature of melting and hydrate decomposition. This result is in accordance with the law of thermodynamic equilibrium. In addition, the increase of pressure also leads to the increase of decomposition heat of gas hydrate. Under the assumption that the structure of hydrate does not change, the increase of decomposition heat can be inferred as more gas entering hydrate and releasing more binding energy. On the other hand, compared with the experimental results of runs 4 and 7, the experiment of run 7 also has two decomposition peaks, but the difference is that the first peak appears melting ice, and the other peak appears hydrate decomposition with a wider shoulder. According to the law of peak-differentiating and fitting, we differentiate and fit the broad peak and get two decomposition peaks at 10.22 \degree C and 14.58 \degree C, respectively. These two temperatures are very close to the corresponding temperature of the two decomposition peaks in the experiment of run 4. And similarly, there should be two kinds of hydrates were formed in the experiment of run 7, namely, CP-CO$_2$ binary mixture hydrate and CO$_2$ hydrate.
of gas hydrate. Under the assumption that the structure of hydrate does not change, the increase of decomposition heat can be inferred as more gas entering hydrate and releasing more binding energy. On the other hand, compared with the experimental results of runs 4 and 7, the experiment of run 7 also has two decomposition peaks, but the difference is more binding energy. According to the law of peak differentiating and fitting, we differentiate and fit the broad peak and get two decomposition peaks at 10.22 °C and 14.58 °C, respectively. These two temperatures are very close to the corresponding temperature of hydrate.

Figure 9. Heat flow and temperature curve at 20.0~−8.8 °C and 0 MPa.

Figure 10. Heat flow and temperature curve at 20.0~−8.8 °C and 3.5 MPa.

Figure 11. Heat flow and temperature curve at 20.0~−8.8 °C and 6.0 MPa.
In fact, the results of Raman measurement on hydrate also proved that the hydrate formed in this system (CP emulsion-CO$_2$) is the coexistence of CP-CO$_2$ hydrate and CO$_2$ hydrate. Figure 14 shows the Raman spectra of the hydrates formed at the initial pressure of 5.0 MPa and temperature of 3.0 °C. On the left side of Figure 14 is the image of the real hydrate formed in the reactor after the complete of the hydrate formation, which is taken by Canon EOS 700 D. The two yellow spots “a” and “b” in the image are the focus of the laser for Raman measurement. The upper and lower images in the middle of Figure 14 are the images of hydrates around “a” and “b” points after being magnified by 500 times, which are captured by the software of Raman spectrometer (Type: Renishaw-In Via Reflex). And on the right side of Figure 14 are the Raman spectra of the hydrates at the spots “a” and “b”. The image on the left side of the figure is the optical micrograph of hydrate before Raman measurement. As seen from the Raman spectrum of the hydrate at the spot “a”, there are four sharp peaks, the two occurring at 1276.3 and 1379.6 cm$^{-1}$ correspond to O=C=O stretch vibrations, indicating that CO$_2$ is included in the hydrate, the other two occurring at 2876.2 and 2983.9 cm$^{-1}$ correspond to the vibration of cyclopentane carbon chain, indicating that CP is also included in the hydrate. In other words, CO$_2$ and CP coexist in the hydrate to form CP-CO$_2$ binary hydrate at the spot “a”. However, from the Raman spectrum of the hydrate at the spot “b”, only two sharp peaks corresponding to O=C=O vibration appear at 1275.7 and 1381.9 cm$^{-1}$, indicating that there is only CO$_2$ in the hydrate at the spot “b”, which is called pure CO$_2$ hydrate. Therefore, the results of Raman analysis prove that the hydrates formed the system are the coexistence of CP-CO$_2$ hydrate and CO$_2$ hydrate. Thus, returning to the discussion of on the results of the experiment.
of runs 4 and 7, it could be concluded that the two decomposition peaks at 10.22 and 14.58 °C represent the decomposition peaks of pure CP-CO₂ binary hydrate and CO₂ hydrate, respectively.

Indeed, we have also considered that these two peaks might be the heat flow peaks generated by the step-by-step decomposition of the same hydrate. For example, for the CP-CO₂ binary hydrate, it is possible that CO₂ and CP molecules successively escape from the cages and absorb heat during the decomposition due to that CP and CO₂ molecules occupy different cages in the hydrate, respectively. Unfortunately, we do not have more scientific evidence to support this view. Although there are successive collapses of different hydrate cages in the molecular dynamics simulation, this kind of phenomenon only occurs in a relatively small time scale. In the macro experimental study, this kind of successive collapses is difficult to verify, let alone in two different hours. In the future work, we will consider whether this possibility is scientific by measuring the single hydrate crystal.
4. Conclusions

In order to reveal the relationship between the microscopic process and the gas consumption of hydrate formation, hydrate composition and formation mechanism of CP-CO$_2$ system was investigated by DSC and Raman measurements on hydrates formed from CP-CO$_2$ emulsion system under different pressures. DSC results show that except the formation and decomposition of ice, there are two peaks of hydrate formation or/and decomposition heat flow, indicating that the two hydrates in hydrate form successively and coexist. At the pressure of 5.0 MPa and above, the endothermic peaks of CP-CO$_2$ binary hydrate and CO$_2$ hydrate decomposition appeared at 10.22 and 14.58 °C, respectively. The Raman spectra also show that the CP-CO$_2$ binary hydrate and the CO$_2$ hydrate coexist in the hydrate phase. The CP-CO$_2$ binary hydrates forms preferentially, and as crystal seeds, they induce the formation of the CO$_2$ hydrates. And the formation process of these two hydrates is the cause of two pressure-drop processes in the whole hydrate formation process. The finding provides a research idea and scientific basis for further improving the gas consumption during the gas hydrate formation in the presence of promoters. It can be possible to increase the total gas consumption by controlling the formation of binary hydrate.

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