Phase Equilibrium and Dissociation Enthalpies for Cyclopentane + Methane Hydrates in NaCl Aqueous Solutions†

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The phase equilibrium data of cyclopentane (CP) + methane hydrates in brine water with NaCl mass fractions of \( w = 0, 0.035, 0.070, \) and 0.100 were measured in the temperature range (284.4 to 301.3) K using a visual high-pressure phase equilibrium apparatus. The dissociation enthalpies of CP + methane hydrates in hot brine water were determined via the Clausius–Clapeyron equation based on these phase equilibrium data. The effect of help gas and salinity on the phase equilibrium of CP hydrate and CP + methane hydrates was studied, respectively. Liquid CP forms hydrates with small-molecule help gas at a temperature region much higher than the quadruple point of pure CP hydrate. Methane partially occupied the small cages of structure II hydrate, which accelerates the nucleation and growth of CP + methane hydrates and increases the stability of the hydrates. The phase equilibrium pressure of CP + methane hydrates increases with the increase in temperature, and it increases linearly with the increase in NaCl concentration in solutions. The higher the temperature, the more remarkable effect the temperature and salinity has on the phase equilibrium pressure. The dissociation enthalpy decreases with the increase in temperature and the NaCl concentration.

Introduction

Gas hydrates (also known as clathrate hydrates) are solid, nonstoichiometric crystalline compounds formed from water and small gas molecules (< 0.9 nm) at high pressures and low temperatures. A huge amount of natural gas hydrates (NGHs) occur naturally within or under permafrost regions and ocean sediments. The most recent estimates of the total amount of methane (STP) trapped in these hydrated gas deposits vary from \( (0.2 \cdot 10^{15} \) to \( 12 \cdot 10^{16} \)) m\(^3\).1 Accordingly, NGHs are considered to be a potential energy resource in the 21st century.2

The methods that have been put forward to destabilize NGHs for gas production include: thermal stimulation,3 depressurization,4 chemical inhibitor injection,5 and replacement of methane by CO\(_2\) hydrate.6 The depressurization method is demonstrated to be highly energy efficient when the NGH deposits are in conjunction with a free-gas zone. However, it will not likely provide a steady, reliable gas production because the dissociation rate of NGHs is controlled by the rate of heat flux from the surrounding media or by the thermal conductivity of the surrounding rock matrix. Thermal stimulation is capable of producing substantial amounts of natural gas, especially when it is combined with the depressurization or chemical injection methods.7 However, thermal stimulation through injection of steam or hot water suffers from significant heat loss during transit of the steam or hot water from the ocean surface to the NGH region. The use of hot brine for dissociation of hydrates was thought be an attractive recovery scheme because it can reduce the dissociation temperature and dissociation enthalpy of NGHs, thus decreasing the requirements of sensible heat and hydrate dissociation energy, as well as heat losses.8

We developed a novel technique to prepare hot brine in situ seafloor for marine NGH exploitation based on hydrate technology.9 In this recovery scheme, a hydrate-forming agent is introduced into an apparatus installed in situ seafloor, in which the hydrate-forming agent forms solid hydrates with seawater spontaneously under submarine high-pressure conditions. The exothermic effect and desalination effect during the formation of hydrate raise the temperature and salinity of the residual seawater, respectively. The hot brine thus prepared is injected into the NGH deposits for NGH dissociation after the solid hydrates float up to the ocean surface by their own buoyancy. The hydrate-forming agent is recovered and recycled at the ocean surface when the solid hydrates float up and decompose into the hydrate-forming agent and water due to the decrease in pressure. The new NGH exploitation process avoids significant heat loss during transiting the heat carrying medium from the ocean surface to the NGH regions in conventional thermal stimulating methods. The hot brine preparation process is to some extent similar to the hydrate desalination process proposed by Max.10 However, the residual brine with relatively high temperature is needed in current technology, while the solid hydrate is needed in Max’s desalination process. Thereby, the hydrate-forming agent and operating parameters are different. The preferred hydrate-forming agent must meet the following requirements: First, the hydrate-forming agent is cheap, nontoxic or low toxic, and insoluble with seawater, and the density of the solid hydrates is lower than that of brine to facilitate its recovery and recycling. Second, the hydrate formation temperature under submarine pressure should be higher than that of methane and as high as possible. Third, the hydrate formation enthalpy should be as high as possible.

Some intermediate/heavy hydrate-forming compounds, such as cyclopentane (CP),11 can form structure II or structure H hydrate at lower pressure and higher temperature conditions than methane by simultaneously use a heavy hydrate former with a
help gas as guest substances. These compounds may be an option for our hydrate-forming agents. The equilibrium pressures of CP + methane hydrate in pure water range from (0.165 to 7.312) MPa in the temperature range (282.15 to 301.90) K. The pressure range is lower than the sub-marinic pressure conditions (ca. (5 to 40) MPa) where common NGH occurs.

Several other studies have reported the dissociation conditions of binary hydrates of CP with different help gases in pure water, such as CH₄, N₂, H₂, CO₂, H₂S, krypton, and difluoromethane. Taylor et al. investigated the hydrate film phase equilibrium pressures of CP in NaCl solutions. The obtained phase equilibrium data of CP + 0.070, and 0.100. The formation/dissociation enthalpies of CP brine water containing the NaCl mass fraction of measured in the temperature range (284.4 to 301.3) K and in the CP/water emulsion system using differential scanning calorimetry. In these experiments, the test fluids were composed of four components (methane, CP, NaCl, and water) and four phases (vapor, liquid hydrocarbon, brine water, and hydrate). The concentration of NaCl in aqueous solution remains constant and is not an independent variable for each given experiment because controlling accuracy of ± 0.1 K. Two Pt100 thermoprobes (JM6051) with ± 0.05 K accuracy were used to measure the temperature at the top (gas phase) and the bottom (liquid phase) of the reactor, respectively. To minimize the perturbation of temperature and pressure in the reactor caused by methane charges, a buffer vessel (1.0 L, 30 MPa) immersed in the temperature-controlled water bath was used to precool and to measure the amount of the supplied gas. The pressure in the reactor was controlled by a proportional-integral-derivative controlled pressure regulated valve (Tescom ER3000) with a pressure-controlling accuracy of ± 0.02 MPa. All pressures in the experimental apparatus were measured using a Danfoss MBS3000 absolute pressure transducer, with a range of (0 to 400) bar and accuracy ± 0.02 MPa. Temperature and pressure data were collected every 6 s using an Agilent model 34970A recording system, which was controlled by the Agilent data acquisition software application BenchLink.

Materials and Preparation of Samples. The methane with the mole fraction of 0.999 was supplied by Guangdong South China Special Gases Technology Institute Ltd., China. CP with the mass fraction of 0.995 was supplied by Chengdu Best Reagent Co., Ltd., China. NaCl with the mass fraction of 0.995 was supplied by Guangzhou Second Chemical Reagent Factory, China. All aqueous solutions were prepared by deionized water which was prepared by an ultrapure water system (Nanjing Ultrapure Water Technology Co., Ltd., China) with a resistivity of 18.25 MΩ·cm.

Procedure. The isothermal pressure search method was employed to determine the hydrate dissociation pressure at a given temperature and a given NaCl concentration in water. The experimental apparatus was washed and rinsed with deionized water, purged with methane three times, and evacuated with a vacuum pump before experimentation. Amounts of 210 mL of NaCl aqueous solution and 25 mL of CP were charged into the reactor under stirring. The buffer vessel was pressurized to the desired pressure with methane. After the temperature of the system was cooled to the predetermined value and remained stabilized by the water bath, the reactor was pressurized with the precooled methane in the buffer vessel to a pressure exceeding the estimated equilibrium pressure by about 1.0 MPa to reduce the induction time. After the hydrate crystal was observed through the glass windows, the pressure was reduced gradually by discharging methane out of the reactor slowly until the hydrate crystals decomposed completely. Then the pressure of the reactor was raised slowly again with the precooled methane in the buffer vessel until a small quantity of hydrate crystal appeared again. When the charging pressure (Pc) of the reactor and the hydrate crystal remained stable for (2 to 3) h, the pressure in the reactor was reduced slowly again to allow the hydrate to decompose. The discharging pressure (Pd) of the reactor was recorded at the moment the hydrate disappeared completely. The above pressure raising and reducing step (or hydrate formation and decomposition process) was repeated until Pd − Pc ≤ 0.05 MPa. At each repetition, the charging pressure Pc was controlled lower than the former repetition, and the discharging pressure Pd was controlled higher than the former repetition. The Pc of the last repetition was taken as the hydrate phase equilibrium pressure at the given temperature and the given NaCl concentration in water.

In these experiments, the test fluids were composed of four components (methane, CP, NaCl, and water) and four phases (vapor, liquid hydrocarbon, brine water, and hydrate). The concentration of NaCl in aqueous solution remains constant and is not an independent variable for each given experiment because...
the hydrate crystal is controlled very small, and it nearly does not affect the NaCl concentration in brine water when the system reaches the phase equilibrium pressure. Thus, according to the Gibbs phase rule, there is only one degree of freedom for the above system at a given NaCl aqueous solution.

Results and Discussion

Effect of Help Gas on Hydrate Equilibrium Conditions of Cyclopentane. The hydrate dissociation conditions of CP and CP + help gas in pure water are shown in Figure 2. Liquid CP form hydrates at a temperature range much higher than the quadruple point of CP, when a help gas, such as CH4, CO2, N2, H2, Kr, or CH2F2 is employed. Cyclopentane can form structure II hydrates without any help gas to fill its small cavities. The pressure of the three-phase equilibrium line of CP (V)—water (LW)—hydrate (H) is lower than atmospheric pressure, and the quadruple point occurred at a temperature of 280.22 K and a pressure of 19.8 kPa.11 The pressures of the three-phase equilibrium line of CP (LHC)—water (LW)—hydrate (H) increase vertically when the temperature is higher than the quadruple point. So there is very little study on hydrate phase equilibrium conditions of pure CP + water at a temperature higher than the quadruple point. At the same time, in a homogeneous nucleation system, CP hydrate could not form until ice occurs and acts as a nucleating agent at a temperature of −38 °C.22,23 Although the supercooling required for heterogeneous nucleation is smaller than that for homogeneous nucleation, CP hydrate is still difficult to form. We had even stirred the CP + water system at 2 °C for 5 days, and there was no CP hydrate generation, though the melting point of CP hydrate is about 7.7 °C at an atmospheric pressure.25 According to the Chen-Guo model,25 cyclopentane forms labile clusters with a number of water molecules surrounding the CP molecule when CP is mixed with water. These clusters, which are the rudiment of a large cage of structure II hydrate, will join together and form basic hydrate when the concentration of these CP — water clusters in water reaches a certain value. The basic hydrate is difficult to grow large enough to create a hydrate nucleus without help gas. However, in the presence of small molecule help gas, these help gases will fill into the linked cavities (small cage of structure II hydrate) of CP basic hydrate, and this increases the stability of the basic hydrate. So the CP basic hydrate can easily grow continuously and form macroscopic hydrate crystals at mild conditions when help gas presents.

The hydrate formation conditions of CP + methane hydrates in water measured in this work are very consistent with the data reported by Sun et al.14 and Tohidi et al.16 The phase equilibrium pressure of CP + methane hydrates in water increase with the increase in temperature. The higher the temperature is, the greater the phase equilibrium pressure increase. The pressures of CP (LHC)—methane (V)—water (LW)—hydrates (H) four-phase equilibrium are in the range (0.48 to 5.694) MPa when the temperatures are in the range (286.67 to 300.02) K. Compared to methane hydrate, the dissociation temperature of CP + methane hydrate is about 20 °C higher than that of methane at the same pressure. Although the equilibrium temperature of CP + methane hydrates is lower than that of CP + Kr and CP + CH2F2 hydrates at the same pressure, CP + methane is considered as a hydrate former in our hydrate-based hot brine preparation because we can take advantage of local materials of seawater and methane recovered from NGHs. The main energy consumption required for the hot brine preparation comes from injecting CP from the ocean surface to the seafloor, as well as separating and recovering of cyclopentane.

Effect of Salinity on Hydrate Equilibrium Conditions of Cyclopentane + Methane. In this work, NaCl aqueous solutions were used as simulated brine water. The four-phase equilibrium data of CP (LHC)—methane (V)—brine water (LBW)—hydrates (H) are listed in Table 1. The equilibrium phase diagrams of LHC—V—LBW—H are displayed in Figure 3. As can be seen from Table 1 and Figure 3a, the phase equilibrium pressures of CP + methane hydrates in brine water increase with the increase

| W     | T   | P   | Z   | Δ Δ diss H m(CH4) / mol (kJ) |
|-------|-----|-----|-----|----------------------------|
| 0.000 | 286.67 | 0.480 | 0.990 | 130.25                     |
| 0.035 | 286.71 | 0.825 | 0.984 | 122.01                     |
| 0.070 | 284.40 | 0.788 | 0.984 | 114.70                     |
| 0.100 | 284.40 | 1.100 | 0.978 | 107.71                     |
| 0.160 | 286.57 | 1.619 | 0.968 | 106.67                     |
| 0.220 | 289.14 | 2.219 | 0.958 | 105.57                     |
| 0.300 | 292.01 | 3.344 | 0.940 | 103.58                     |
| 0.500 | 295.67 | 5.788 | 0.907 | 99.95                      |
| 0.700 | 299.33 | 8.800 | 0.878 | 92.39                      |
| 1.000 | 302.83 | 11.888 | 0.861 | 100.33                     |
| 1.500 | 305.75 | 16.344 | 0.848 | 93.44                      |
hydrates in hot brine takes an important role in our hydrate-water increase linearly with the increase in NaCl mass fraction phase equilibrium pressures of CP

mass fraction increases at a high temperature. For the solutions with the NaCl on the phase equilibrium pressures of CP

from ca. (286 to 301) K. This indicates that the effect of salinity CP

in temperature. The increasing rates of the phase equilibrium pressures increase with the increase in temperature. NaCl in solutions has an obvious inhibition effect on the formation of CP + methane hydrates. The phase equilibrium pressures of CP + methane hydrates in brine water were significantly higher than that in pure water. As can be seen from Figure 3b, the phase equilibrium pressures of CP + methane hydrates in brine water increase linearly with the increase in NaCl mass fraction (salinity) in solutions. The slopes of $P - 100\cdot w$ lines increase with the increase in temperature. The slopes of $P - 100\cdot w$ lines increase from 0.1113 to 0.8880 when the temperatures increase from ca. (286 to 301) K. This indicates that the effect of salinity on the phase equilibrium pressures of CP + methane hydrates increases at a high temperature. For the solutions with the NaCl mass fraction $w = 0.035$ and 0.070, the phase equilibrium pressures of CP + methane hydrate at 301.31 K are (10.513 and 11.888) MPa, respectively. These pressures are lower than the seafloor pressure where common oceanic NGH formation occurs, (5 to 40) MPa.

**Dissociation Enthalpy of Cyclopentane + Methane Hydrates.** The formation/dissociation heat of CP + methane hydrates in hot brine takes an important role in our hydrate-based hot brine preparation, and these dissociation enthalpy data are required to develop production schemes. Dissociation enthalpies of gas hydrates are usually obtained by direct calorimetric measurement and indirect determination via the Clapeyron or Clausius–Clapeyron equation by differentiation of phase equilibrium pressure–temperature data. The formation and dissociation of CP + methane hydrates can be represented by the following equation

$$
\text{CH}_4 + x\text{CP} + n\text{H}_2\text{O} \leftrightarrow \text{CH}_4 \cdot x\text{CP} \cdot n\text{H}_2\text{O} \quad (1)
$$

where $x$ and $n$ are the mole number of CP and H$_2$O in hydrate crystalline, respectively.

To a first-order approximation, the hydrate dissociation enthalpy above 273.15 K can be determined from the univariant slope of the phase equilibrium line of ln(P/MPa) vs 1000 K/T using the Clausius–Clapeyron equation

$$
\frac{d\ln P}{d(1/T)} = 
-\frac{-\Delta_{\text{diss}}^\circ H_{\text{m(CH}_4)}^{\text{ch}}}{ZR} 
$$

(2)

where $P$ is the pressure; $T$ is the temperature; $Z$ is the compression factor of the guest gas (CH$_4$) at each equilibrium $P$–$T$ condition, and it can be calculated by the Soave–Redlich–Kwong (SRK) equation; $R$ is the universal gas constant ($R = 8.31451$ J·mol$^{-1}$·K$^{-1}$); and $\Delta_{\text{diss}}^\circ H_{\text{m(CH}_4)}^{\text{ch}}$ is the molar dissociation enthalpy of CP + methane hydrates calculated by methane.

The $(P, T)$ data in Table 1 are plotted as ln(P/MPa) vs 1000 K/T in Figure 4. As can be seen from Figure 4, ln(P/MPa) vs 1000 K/T exhibits a very good linear relation under all NaCl concentrations. The slopes $k$ in Figure 4 increase linearly with the increase in NaCl concentration in aqueous solution, which can be described by the linear equation of $k = 25656.7 w - 15816.657$ with a linear correlation coefficient $r = 0.9976$.

The dissociation enthalpies of CP + methane hydrates in hot brine calculated by eq 2 are listed in Table 1 and displayed in Figure 5. As can be seen from Figure 5, $\Delta_{\text{diss}}^\circ H_{\text{m(CH}_4)}^{\text{ch}}$ decreases with the increase in temperature and the increase in phase
nucleation and growth of CP + methane hydrates and increases its stability. NaCl in solutions has an obvious inhibition effect on the formation of CP + methane hydrate. The phase equilibrium pressure of CP + methane hydrates increases with the increase in temperature, and it increases linearly with the increase in NaCl concentration. The higher the temperature, the more remarkable effect the temperature and salinity have on the phase equilibrium pressure. The dissociation enthalpies of CP + methane hydrates in hot brine water were determined via Clausius–Clapeyron equation based on the measured phase equilibrium data. The dissociation enthalpy decreases with the increase in temperature and the increase in NaCl concentration because the increase in temperature and the salinity of solutions increase the phase equilibrium pressure, which caused the decrease of \( Z \) and the increase of the fractional occupancy of cages by methane.

### Conclusions

The phase equilibrium data of CP + methane hydrates were obtained in brine water with the NaCl mass fraction of \( w = 0, 0.035, 0.070, \) and \( 0.100, \) in the temperature range from (284.4 to 301.3) K, using a visual high-pressure phase equilibrium apparatus. Liquid CP forms hydrate at a temperature region much higher than the quadruple point of pure CP hydrate when small molecule help gas presents. Methane partially occupies the small cages of structure II hydrate, which accelerates the

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**Figure 5.** Dissociation enthalpy of cyclopentane + methane hydrates versus temperature (a) and pressure (b) in different NaCl solutions. \( \Delta H_{\text{diss}} \) at 0.000; \( \bullet \), \( w = 0.035 \); \( \Delta \), \( w = 0.070 \); \( \nabla \), \( w = 0.100 \); \( \Delta H_{\text{diss}} \) of CH4 dissociation enthalpy calculated by CH4.
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