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Crystal Structure of Mixed Hydrates and Cage Occupancy of Guest Species

YUUKI MATSUMOTO

MARCH 2014

Crystal Structure of Mixed Hydrates and Cage Occupancy of Guest Species

A dissertation submitted to THE GRADUATE SCHOOL OF ENGINEERING SCIENCE OSAKA UNIVERSITY in partial fulfillment of the requirements for the degree of DOCTOR OF PHILOSOPHY IN ENGINEERING

BY

YUUKI MATSUMOTO

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Abstract

Clathrate hydrate is suggested to be utilized as gas storage and transportation media and refrigerants because of its dense storage capacity and high dissociation enthalpy, respectively. Generally, clathrate hydrate is thermodynamically stable at high pressures and/or low temperatures, and its stability largely depends on guest species. One of the most efficient ways to moderate the thermodynamic stability conditions of clathrate hydrate is addition of relatively large guest species called thermodynamic promoters to form a mixed hydrate of a target gas with a thermodynamic promoter. Compared to simple hydrate systems, the phase behavior and the cage occupancies of guest species in mixed hydrate systems are much more complicated. In this thesis, thermodynamic promoters were explored toward moderating the thermodynamic stability conditions of a simple hydrate composed of a target gas such as hydrogen, methane, or carbon dioxide. Thermodynamic stability conditions and crystal structure of mixed hydrates were also investigated by means of phase equilibrium measurement and powder X-ray diffractometry. Simultaneously, cage occupancies of guest species in mixed hydrates were clarified by means of Raman spectroscopy. Mixed hydrate systems investigated in the present study were classified according to the cage occupancy; competitive cage occupancy (Competitive-type, Chapter II) and compartmental cage occupancy (Compartmental-type, Chapter III).

Competitive-type mixed hydrates, where two or more kinds of guest species competitively occupy a certain cage, are summarized in **Chapter II**. Isothermal phase equilibrium relations (pressure-composition) of mixed hydrate systems exhibited some interesting phase behavior such as structural phase transition and azeotropic-like behavior. According to structural phase transition, mixed hydrates construct the most thermodynamically stable crystal structure depending on the cage occupancy of guest species at a given pressure, temperature, and composition. Azeotropic-like behavior indicates that, at a local pressure minimum, guest species compartmentally occupy the appropriate cages in spite of competitive-type mixed hydrate.

Compartmental-type mixed hydrates, where small and large guest species occupy small and large cages selectively, are summarized in **Chapter III**. In general, the thermodynamic stability conditions and crystal structures of mixed hydrates mainly depend on the large guest species. The present results elucidate that the cage occupancy of large guest species in mixed hydrate depends also on small guest species. An unexpected high-stability of the compartmental-type mixed hydrate system at high pressures and high temperatures was discovered, which suggests the competitive large-cage occupancy by large guest species and small guest species.

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Chapter I

General Introduction

I.1. Properties of Clathrate Hydrate

Clathrate hydrate (hereafter simply called "hydrate") is a solid crystalline, which has an icelike appearance. Hydrate is necessarily composed of host species and guest species, where the former is water (H₂O) molecules and the latter is generally small molecules such as light hydrocarbons and noble gases. Several host molecules construct cage structure (hereafter called "cage") and a cage can normally enclathrate one guest molecule. Generally, a cage enclathrating a guest molecule is more stable than a vacant cage. There are several kinds of hydrate cages. Typical cages are shown in Figure I-1. For instance, 5¹²-cage (S-cage) is a dodecahedron, which is constructed from twelve (¹²) pentagons (5). The other cages (S'-cage, M-cage, L-cage, and U-cage) are summarized in Table I-1. S-cage, S'-cage, and L-cage are nearly spheral but M-cage and Ucage are oval. The other properties of the typical cages are also summarized in Table I-1. Some of them are combined to form specific hydrate structures as shown in Figure I-2. The unit cell of the structure-I (s-I) hydrate consists of two S-cages and six M-cages. In the same manner, that of the structure-II (s-II) hydrate is formed with sixteen S-cages and eight L-cages. The unit cell of structure-H (s-H) hydrate is composed of three S-cages, two S'-cages, and one U-cage. The properties of s-I, s-II, and s-H hydrates are summarized in Table I-2 accompanied with those of ice-Ih. Simple hydrate is formed with binary mixture of host species and single guest species. On the other hand, mixed hydrate is formed with more than ternary mixture that is host species and two or more kinds of guest species. Simple hydrates normally belong to s-I or s-II hydrate, and mixed hydrates belong to s-I, s-II, s-H, or the other structures. Crystal structures and thermodynamic stabilities of hydrates depend on the thermodynamic conditions (pressure, temperature, composition, etc.) and the properties of guest species (size, shape, molar volume, molar weight, solubility, polarity, etc.) [1,2]. A lot of guest species have been reported until now. The guest species investigated in the present study are shown in Figure I-3 in the ascending order of van der Waals diameter. The guest species can be categorized as; (a) occupying S-cages and L-cages of s-II hydrate, (b) occupying S-cages and M-cages of s-I hydrate, (c) occupying M-cages of s-I hydrate, (d) occupying L-cages of s-II hydrate, (e) occupying L-cages of s-II hydrate with help gas, and (f) occupying U-cages of s-H hydrate with help gas ("help gas" is explained below). The guest species categorized as (a) are very small and form s-II hydrates because s-II hydrates have a larger ratio of small cages to large cages than that of s-I hydrates. Some of guest species can multiply occupy Lcages. For instance, hydrogen (H₂) molecules occupy an L-cage doubly, triply, or quadruply [3]. The guest species in (b) such as methane (CH₄) and xenon (Xe) are slightly larger than those of (a), and they occupy both S-cages and M-cages of s-I hydrates. The guest species in (c) are larger than the cavity of S-cage. Therefore, they occupy only M-cages of s-I hydrates. At high pressures, however, some of them such as ethane (C_2H_6) [4], ethylene (C_2H_4) [5], cyclopropane (c- C_3H_6) [6], and tetrafluoromethane (CF₄) [7] are compressed and forced to occupy S-cages. The guest species in (d) are larger than the cavity of M-cage. Therefore, they occupy only L-cages of s-II hydrates. As exceptions, sulfur hexafluoride [8] and propane (C_3H_8) [9], which are in (d), are compressed and forced to occupy M-cage of s-I hydrates at high pressures. The guest species in (e) are slightly large compared to the cavity of L-cage to occupy. Therefore, they cannot form s-II hydrate solely. When the small guest species such as CH₄ are added, large guest species occupy L-cages of s-II hydrates with the help of small guest species occupying S-cages. This is why the small guest species is called "help gas". The guest species in (f) are larger than the cavity of L-cage, so they occupy U-cages of s-H hydrates with the support of help gas occupying S-cages and S'-cages. In the case of s-H hydrates, the ratio of small cages to large cages is larger than that of s-I and s-II hydrates, and S'cage and U-cage have geometrically unstable faces (squares and hexagons). Therefore, stabilization of cages by help gas occupying S-cages and S'-cages is necessary to form s-H hydrates [10].



Figure I-1. Schematic illustration of typical hydrate cages.

Name	S-cage	S'-cage	M-cage	L-cage	U-cage
Туре	512	435663	51262	51264	51268
Kind of polyhedron	dodecahedron	dodecahedron	tetrakaideca- hedron	hexakaideca- hedron	icosahedron
Coordination number	20	20	24	28	36

Table I-1. Summary on properties of typical hydrate cages.



Figure I-2. Schematic illustration of crystal structures of hydrates.

Solid phase	s-I hydrate		s-II hydrate		s-H hydrate		ice-Ih	
Cage	S	М	S	L	S	S'	U	-
Number of cages	2	6	16	8	3	2	1	-
Cage diameter d_1 / nm	0.795	0.860	0.782	0.946	0.782	0.812	1.142	-
Cavity diameter d_2 / nm ^a	0.51	0.59	0.50	0.67	0.49 ^b	0.51 ^b	0.86 ^b	-
Number of water molecules	4	-6	1.	36		34		-
Crystal type	Cubic		Cu	bic	H	Iexagon	al	Hexagonal
Space group	Pm3n		Fd	3m		P6/mmn	ı	P6 ₃ /mmc
Lattice constant a / nm	1.20 ^c		1.7	73 ^c		1.22^{d}		0.451842^{e}
Lattice constant c / nm -			-		1.01 ^d		0.735556 ^e	

Table I-2. Summary on unit cells of hydrates and ice-Ih.

^a The cavity diameters are obtained from the cavity diameter minus the diameter of water (0.28 nm).

^b From the atomic coordinates measured using single crystal X-ray diffraction on s-H hydrates [11].

^{*c*} From ref. [12].

^{*d*} From ref. [13].

^{*e*} From ref. [14] (at T = 250 K).



Figure I-3. Relation among size of guest species, hydrate structure, and cage occupancy investigated in the present study.

I.2. History and Applications of Clathrate Hydrate

Davy discovered water-chlorine (Cl₂) crystalline compounds, which were later identified as hydrates, in 1811 [15]. Faraday and Davy clarified that the mole ratio of Cl₂ to water was 0.1 (Cl₂·10H₂O) [16]. Pauling and Marsh found out that the Cl₂ crystals were composed of host water molecules and guest Cl₂ molecules by means of X-ray diffractometry, that is, the crystals were "clathrate hydrates" [17]. After their studies, the structure models of s-I and s-II hydrates were indicated with a X-ray diffractometer [18]. In 1987, s-H hydrate was discovered by means of ²H and ¹²⁹Xe NMR spectroscopy and neutron powder diffractometry by Ripmeester *et al.* [10].

In nature, large amounts of natural gas hydrate exist in permafrost and marine sediments as shown in **Figure I-4** [19]. Natural gas hydrate as well as shale gas, tight sand gas, and coal bed methane is non-conventional energy source. Japan is one of the energy-poor countries. However, there are large amounts of natural gas hydrate in deep sea around Japan. The estimated deposit of natural gas hydrate around Japan is approximately a hundred times of the annual consumption of natural gas in Japan [20]. Exploitation of the natural gas hydrate is significantly important for the energy self-sufficiency and the price negotiation of imported natural gas.



- O Natural gas hydrate fields under the deep ocean floor
- Natural gas hydrate fields in the permafrost regions

Figure I-4. Distribution of the natural gas hydrate field in the world [19].

In U.S., hydrate has been considered as a nuisance because it forms in the petroleum pipelines and blocks them [21]. This blockage accident occasionally happened. Therefore, main research topic on hydrate has been to explore inhibitors of the hydrate formation. At first, salt and alcohol were investigated as thermodynamic inhibitors that make hydrate thermodynamically unstable as shown in **Figure I-5**. Then, low molecular weight polymers were investigated as kinetic inhibitors that delay hydrate formation. Recently, surfactants have been investigated as anti-agglomerants that inhibit aggregation of hydrate particles.



Figure I-5. Phase equilibrium relation for CH₄ hydrate systems with and without additives.

As mentioned before, hydrate can enclathrate large amount of gas. For example, an ideal CH₄ hydrate has approximately 175 times of its volume of CH₄ at standard conditions. This property can give us a potential as the alternative medium for gas storage, transportation, and isolation to gas compression, gas liquefaction, or gas storage in metal hydrides. However, hydrate formation and stabilization requires high pressures and/or low temperatures, which is one of the most severe

problems to be solved. The most effective and simplest solution is an addition of thermodynamic promoter such as cyclopentane (c-C₅H₁₀) and tetrahydrofuran (THF), which is larger than the gas molecule and occupies large cages of hydrates. By adding thermodynamic promoters, mixed hydrate forms at more moderate thermodynamic conditions than simple gas hydrate. For example, the thermodynamic stability regions of CH₄+additive mixed hydrates are larger than that of simple CH₄ hydrate as shown in **Figure I-5**. A negative aspect of an addition of thermodynamic promoter is the decrease of the amount of gas enclathration. For the applications of hydrate for gas storage, transportation, and isolation, it is better to achieve both the moderation of hydrate formation condition and the retention of gas enclathration amount.

Hydrogen Hydrate

Originally, it had not seemed that H₂ can form hydrate because of its tiny size. In 1999, H₂ hydrate was firstly reported in the phase diagram for H₂+water system as shown in Figure I-6 by Dyadin et al. [27]. H₂ molecules form s-II hydrates and occupy S-cages singly and L-cages doubly, triply, and quadruply [28]. According to molecular simulation, H₂ molecules can occupy S-cage doubly [29]. However, there is no experimental evidence so far. When it was assumed that H₂ molecules occupy all the S-cages and L-cages singly and quadruply, respectively, the ideal storage capacity of H₂ hydrate is 3.8 wt% of H₂. However, simple H₂ hydrate requires extremely severe thermodynamic conditions for its formation and stabilization, which is the reason why H₂ hydrate is not applied practically yet. In the case of H₂ hydrate, especially, the effect of thermodynamic promoters is significant, compared to the other hydrates such as CH₄ hydrate and carbon dioxide (CO₂) hydrate. For example, s-II H₂+THF hydrate forms at p = 10 MPa and T = 280 K, on the other hand, simple H₂ hydrate forms at p = 150 MPa and T = 270 K [30,31]. In addition, the thermodynamic stability of the H₂+THF hydrate depends on the THF concentrations as shown in Figure I-7 [32]. Recently, some kinds of quaternary ammonium salts have been reported as new thermodynamic promoters forming semi-clathrate hydrate. H₂+tetra-n-butyl ammonium bromide semi-clathrate hydrate has high stability as well as H₂+THF hydrate [31]. In the case of H₂+THF hydrate, however, the storage amount of H₂ decreases down to 1 wt% because THF molecules occupy all of the L-cages of s-II hydrates and there is no L-cage available for H₂ enclathration. Recently, it was reported that H₂ is allowed to occupy L-cage multiply in H₂+THF hydrate by decreasing the THF concentration and using special approach, which is called "tuning effect" [33,34]. Ideally, the amount of H₂ in the tuned H₂+THF hydrate achieves up to 3.8 wt% and the tuned hydrate forms outside the stable region of simple H₂ hydrate. The tuned hydrate is at the metastable state. Therefore, the tuning effect significantly depends on all of the experimental conditions and procedures.



Figure I-6. Phase diagram for H₂+water system [27].



Figure I-7. Phase diagram for H₂+THF+water system [31,32].

Methane Hydrate

CH₄ hydrate has been investigated for a long time. Because CH₄ hydrate is flammable (as shown in **Figure I-8**) in spite of its ice-like appearance, it is called "flammable ice". As mentioned above, natural gas hydrate exists in permafrost field and marine sediment [19]. Complexly, natural gas hydrates include not only CH₄ but also C_2H_6 , C_3H_8 , *n*-butane, *i*-butane, and other components. It is important to know the thermodynamic stability of many kinds of natural gas hydrate for its exploitation. On the other hand, it is also suggested that CH₄ hydrate is applied for natural gas storage and transportation as shown in **Figure I-9**. Nowadays, natural gas is transported as liquefied natural gas. However, extremely low-temperature is required for its liquefaction (T = 110 K at p = 0.1 MPa). Hydrate is allowed to store and transport natural gases at a condition more moderate than that of liquefied natural gas. Furthermore, it is reported that the dissociation rate of CH₄ hydrate is reduced at a certain temperature region at atmospheric pressure as shown in **Figure I-10** [35], so-called "self-preservation phenomenon". It is much more desirable to use self-preserved CH₄ hydrate is superficially stable at p = 0.1 MPa and T = 263 K.



Figure I-8. Photo of burning CH₄ hydrate.



Figure I-9. Schematic illustration of storage and transportation of natural gas using hydrates.



Figure I-10. Dissociation rate of CH₄ hydrate depending on temperature at atmospheric pressure [35].

Carbon Dioxide Hydrate

 CO_2 is one of the materials which cause global warming. Avoiding this worldwide problem, it is necessary to reduce CO_2 emission and to promote CO_2 isolation (called "carbon capture and storage (CCS)") and CO_2 utilization. As a method for CO_2 isolation, it is suggested to use CO_2 hydrate. Just leading CO_2 to sea bottom deeper than 6000 m, CO_2 is liquefied at the bottom and at the interface between sea water and liquefied CO_2 , CO_2 hydrate is formed as shown in **Figure I-11** [36]. As a result, liquefied CO_2 can be isolated very densely beneath CO_2 hydrate film. Furthermore, it is suggested that CH₄ enclathrated in natural gas hydrate is replaced with CO₂ by supplying CO₂ into the sand layer including natural gas hydrates as shown in Figure I-12 [37]. This process is more valuable because CO₂ isolation and CH₄ collection can be achieved simultaneously. As CO₂ utilization, CO₂ is suggested to be used as a refrigerant [38]. Hydrate has some positive aspects for refrigerant. Because the dissociation enthalpy of the gas hydrate is higher than the fusion enthalpy of ice, the hydrate is often employed as thermal energy storage and hydrate-based refrigeration systems [39]. Generally, gas hydrates have the dissociation temperature higher than ice at some pressure, and the dissociation temperature of hydrates can be controlled by changing pressure or adding the other guest species, whereas it is so hard and not effective to control the fusion temperature of ice with pressurization or depressurization. The usage of hydrate slurries can be applied for not only batch cooling system but also continuous one. CO₂ hydrate is more appropriate for refrigerant than hydrocarbon hydrates because the former has the higher dissociation enthalpy than the latters. However, high pressures and low temperatures are still required to form CO₂ hydrates. Thus, it is proposed to moderate the conditions of hydrate formation by use of thermodynamic promoters.



Figure I-11. CO₂ isolation process using hydrates [36].



Figure I-12. CH₄/CO₂ replace process using hydrates [37].

Hydrofluorocarbon Hydrate

Hydrofluorocarbons (HFCs) have been used as refrigerants instead of chlorofluorocarbons and hydrochlorofluorocarbons because chloride compounds destroy the ozone layer. HFCs still significantly promote the global warming as similar to CO₂. However, HFC hydrates form under conditions much more moderate than CO₂ hydrate as shown in **Figure I-13**. HFCs generally have global warming potential higher than CO₂, and some of HFCs are flammable. Therefore, it is desirable to use these hydrates as refrigerant in completely closed system.



Figure I-13. Phase diagram for simple CO₂ and HFC hydrates.

I.3. Research Purposes of the Present Studies

As mentioned in the previous section, many kinds of utilizations using hydrates have been proposed. However, some problems to be solved still remain. For H₂ storage, it should be satisfied to achieve both the moderation of thermodynamic conditions of H₂ hydrate and the retention of H₂ enclathration amount. Therefore, the mechanism of tuning effect should be clarified to handle it. Or the alternative storage method using hydrate (additive or procedure) should be established. For natural gas storage and transportation, the practical hydrate system should be investigated for feasible natural gas storage and transportation process. For refrigerants, it is important to form CO₂ hydrate or HFC hydrate at a pressure as low as possible for safe, low-cost, and feasible cooling process. What is the comprehensively important for hydrate utilization is to discover remarkable thermodynamic promoters and to use mixed hydrates composed of a target gas and the thermodynamic promoter. Compared to simple hydrate systems, mixed hydrate systems exhibit complicated phase behavior and cage occupancies of guest species. In the present study, small guest

species (H₂, CH₄, CO₂, etc.) and many kinds of the candidates of thermodynamic promoter were adopted. Thermodynamic stabilities of many mixed hydrates were clarified by means of phase equilibrium measurement. The crystal structures of unclarified mixed hydrates were identified by means of powder X-ray diffractometry (PXRD). In addition, cage occupancies of guest species in mixed hydrates were analyzed by means of Raman spectroscopy. Taking into consideration all of the experimental results, phase behavior and structural transitions depending on cage occupancies of guest species in mixed hydrates were comprehensively discussed. According to the cage occupancies, the mixed hydrate systems investigated in this thesis were categorized into two types; competitive-type mixed hydrates are summarized in **Chapter II** and compartmental-type mixed hydrates are in **Chapter III**.

I.4. Nomenclatures

a	Lattice constant [m]
с	Lattice constant [m]
d_1	Cage diameter [m]
d_2	Cavity diameter [m]
р	Pressure [Pa]
Т	Temperature [K]
<i>W</i> CH3OH	Mass fraction of methanol in aqueous solution [-]
WNaCl	Mass fraction of sodium chloride in aqueous solution [-]
XTHF	Mole fraction of THF in aqueous solution [-]
G	Gas phase
Н	Hydrate phase
L_1	Aqueous phase
S_1	Ice phase

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Chapter II

Competitive-Type Mixed Hydrates

Section 1

In Situ Raman Spectra of Hydrogen in Large Cages of Hydrogen+Tetrahydrofuran Mixed Hydrates

II.1.1. Abstract

Large-cage occupancy of H₂ in the H₂+THF hydrates has ever been reported only in the *ex* situ spectroscopic measurements with the quenched hydrate sample. In this section, to verify the proof-of-concept for the large-cage occupancy of H₂ competitive with THF, *in situ* Raman spectra of H₂ in the H₂+THF mixed hydrates were detected at p = 74.3 MPa and T = 265 K without the quenching procedure. The Raman spectra revealed that H₂ molecules occupy a part of large cages of the H₂+THF mixed hydrates in a thermodynamically unstable region of simple H₂ hydrate.

II.1.2. Introduction

H₂ and water form s-II hydrate [1,2]. The unit cell of s-II clathrate hydrate consists of sixteen S-cages and eight L-cages, which are singly and multiply occupied by H₂ molecules, respectively [3]. Simple H₂ hydrates potentially offer a clean H₂ storage medium. However, extremely high pressures and/or low temperatures are required for simple H₂ hydrate formation [1–3]. The addition of THF makes H₂+THF mixed hydrates at pressures lower than the equilibrium ones of simple H₂ hydrates [4–9]. In the H₂+THF mixed hydrates prepared with compressed H₂ and THF aqueous solutions, H₂ molecules occupy only S-cages, while THF molecules occupy L-cages [10]. That is, the cage occupancy of H₂+THF mixed hydrates are prepared with compressed H₂ and solid mixtures of THF and ice, H₂ molecules metastably occupy not only S-cages but also a part of L-cages [11,12]. The phenomenon has been called the tuning effect [5]. In this case, H₂ storage amount ideally increases to the similar level as that of simple H₂ hydrates. Until now, competitive L-cage occupancies of H₂ and THF have been investigated by means of *ex situ* Raman spectroscopy at atmospheric pressure (p = 0.1 MPa) and liquefied nitrogen (LN₂) temperature (T = 77 K) [11,12]. The hydrate samples

were taken out from high-pressure cells after they were quenched by LN_2 . Therefore, some researchers have inferred that the peaks derived from H_2 molecules in L-cages were detected as a result of simple H_2 hydrates generation during the quenching process.

In this section, H₂+THF mixed hydrates were prepared and analyzed in simple H₂ hydrate unstable region (p = 74.3 MPa and T = 265 K).

II.1.3. Experimental

Materials

The materials used in the present study are summarized in **Table II-1-1**. All of them were used without further purification.

 Table II-1-1. Information on the materials used in the present study.

Material name	Source	Mole fraction purity		
H ₂	Neriki Gas Co., Ltd.	> 0.999999		
THF	Merck, Ltd.	> 0.999 (Main impurity: water)		
Distilled water	Wako Pure Chemicals Ind., Ltd.	> 0.9999		

Apparatus

A schematic illustration of the high-pressure optical cell used in the present study is shown in **Figure II-1-1**. The cell has a pair of titanium-free sapphire windows. A laser Raman confocal microprobe spectrophotometer (Jasco, NRS-1000) with a multichannel charge-coupled device (CCD) detector was used accompanied with an argon ion laser (wavelength: 514.5 nm, laser spot diameter: 2 μ m). The spectral resolution of the spectrometer was 0.7 cm⁻¹. The system temperature was measured using a RTD thermistor (Eutech, Temp 6, reproducibility: 0.2 K). The system pressure was measured by a pressure gauge (Valcom, VPRH, maximum uncertainty: 0.2 MPa).



Figure II-1-1. Schematic illustration of the experimental setup and procedure.

Procedure

In the present study, to exclude both effects of the sample quenching and the fluid H₂ in the Raman analysis, I used a different experimental method from the previous reports [10,12]. A schematic illustration of the experimental setup and procedure is shown in Figure II-1. A small droplet of pure water was directly dropped on a sapphire window of the high-pressure optical cell. It was placed in a temperature-controlled room at T = 263 K, and the droplet was frozen. Then the window with ice was set in the main unit of high-pressure cell, and THF was enclosed within a high-pressure pipe line between the valve and the high-pressure optical cell. Once the valve was opened, THF was sprayed on the ice surface by compressed H₂. At this time, H₂ and THF molecules simultaneously supplied to the ice surface result in the hydrate formation. The approximate amount of the enclosed THF corresponded to the stoichiometric concentration of the simple THF hydrate because the hydrate formation was unable to be detected under a less THF concentration. Though it is difficult to comprehend the exact amount of THF reacted with ice, the THF concentration (on an H₂ free basis) in the H₂+THF mixed hydrate is at least less than the stoichiometric concentration of the simple THF hydrate. The system temperature was controlled at T = 265 K. The hydrate samples were analyzed through a sapphire window by means of *in situ* Raman spectroscopy at p = 74.3 MPa and T = 265 K. The argon ion laser was irradiated to the hydrate crystal and the backscatter was taken in through the same lens. To maximally inhibit the overlap of fluid H₂ on the Raman spectrum, that is, to improve the spatial resolution in the confocal Raman arrangement, the minimal sizes of a slit and an aperture were used. As a compensation of the improved spatial resolution, detected peak intensities are generally sacrificed.

II.1.4. Results and Discussion

Raman peaks corresponding to the intramolecular H–H stretching vibration mode of H₂ in Lcage are strongly affected by surrounding temperature [2]. It has been reported that the peak derived from single H₂ molecule in S-cages (hereafter, peak(S)) is detected at $\Delta v = 4132$ cm⁻¹ and the peaks derived from multiple H₂ molecules in L-cages (peaks(L)) are detected at $\Delta \nu = 4144$ cm⁻¹ and 4152 cm^{-1} at T = 77 K [13]. Note that the peak widths and positions are different from those recorded at an extremely low temperature and the peak(S) is broad and seems to be single at ambient temperature. Figure II-1-2 shows the Raman spectra of the H–H stretching vibration mode of H₂ molecules in the H₂+THF mixed hydrate and fluid H₂ phases. Figure II-1-2(a) indicates the Raman spectrum of H₂ in the H₂+THF mixed hydrates prepared with the present experimental method. Figure II-1-2(a) also involves the slight contribution of the fluid H₂. It was very difficult, even with the confocal Raman arrangement, to completely exclude the effect of H₂ in fluid phase. Figure **II-1-2(b)** is the spectrum of the H_2 +THF mixed hydrate prepared from compressed H_2 and THF aqueous solutions. In this case, no H₂ molecule occupies L-cage even at low THF concentrations [10]. The peak(S) in Figure II-1-2(a) has a shoulder at the high wavenumber side, while it does not appear in Figure II-1-2(b). In order to elucidate that the peaks(L) exist in Figure II-1-2(a), a difference spectrum between Figure II-1-2(a) and Figure II-1-2{(b)+(c)} is shown in Figure II-1-3(a). In this calculation, Figure II-1-2(b) and Figure II-1-2(c) were normalized by the intensities of the peaks detected at $\Delta v = 4132$ cm⁻¹ and 4159 cm⁻¹, respectively. Figure II-1-3(a) indicates two peaks assuredly at $\Delta v = 4140-4160$ cm⁻¹. They were fitted by use of Voigt function with the base line correction. The deconvoluted peaks are shown in the dotted ($\Delta v = 4144$ cm⁻¹) and dashed ($\Delta v = 4152 \text{ cm}^{-1}$) lines. They are in good agreement with the peaks(L) of simple H₂ hydrates [13] as shown in **Figure II-1-3(b)**. This result reveals that H_2 molecules occupy a part of L-cages in the H₂+THF mixed hydrates prepared with the present experimental method. In other words, the competitive L-cage occupancy of H₂ with THF in H₂+THF mixed hydrates surely occurs, and it is not always derived from the simple H₂ hydrate formation. The lower L-cage occupancy of H₂ than previous reports [11,12] results from the present experimental procedure to verify the proof-ofconcept, that is, to exclude the effect of fluid H₂ in Raman analysis. Optimizing a preparation procedure and condition would make improvements.



Figure II-1-2. Raman spectra corresponding to the H–H stretching vibration mode of H₂ molecules in the H₂+THF mixed hydrates and fluid phase. Spectrum (a) partially includes fluid H₂.



Figure II-1-3. (a) Difference spectrum between **Figure II-1-2(a)** and **Figure II-1-2{(b)+(c)}**, and (b) the Raman spectrum of H₂ in the simple H₂ hydrate [13].
II.1.5. Conclusions

The competitive L-cage occupancy of H_2 molecules in H_2 +THF mixed hydrates has been *in situ* observed in a thermodynamically unstable region of the simple H_2 hydrate. The additional Raman peaks, other than the peak derived from single H_2 molecule in S-cage, reveal the competitive L-cage occupancy of H_2 with THF (so-called tuning effect), while the cage occupancy of the H_2 +THF mixed hydrates belongs to the compartmental-type on the thermodynamic stability.

II.1.6. Nomenclatures

ν	Wavenumber [m ⁻¹]
р	Pressure [Pa]
Т	Temperature [K]
x_{THF}	Mole fraction of THF in aqueous solution [-]

II.1.7. References

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Section 2

Thermodynamic Stabilities of Hydrogen+Methane Binary Gas Hydrates

II.2.1. Abstract

When H_2 is mixed with small amounts of CH_4 , the conditions required for clathrate hydrate formation can be significantly reduced when compared to that of simple H_2 hydrate. With growing demand for CH_4 as a commercially viable source of energy, H_2+CH_4 binary hydrates may be more appealing than extensively studied H_2+THF hydrates from an energy density standpoint. Using Raman spectroscopic and PXRD measurements, I show that hydrate structure and storage capacities of H_2+CH_4 mixed hydrates are largely dependent on the composition of the initial gas mixture, total system pressure, and formation period. In some cases, H_2+CH_4 hydrate kinetically forms s-I first, even though the thermodynamically stable phase is s-II.

II.2.2. Introduction

As the amount of easily accessible petroleum-based energy resources continues to decline, the need for alternative energy materials is crucial. Of the recently proposed alternatives, two of the more promising include the use of H_2 and natural gas (e.g. CH₄). Examining the current technologies, many ways exist to store and transport these gases such as, simple compression, liquefaction, metal hydrides, and metal organic frameworks [1]. Alternatively, with the ability to concentrate gases over 175 times their volume at standard conditions, clathrate hydrates, such as those occurring naturally in sediments under the ocean floor and in permafrost regions, have also been considered a promising energy storage medium [2–5]. However, whereas the structure and thermodynamics of natural gas hydrates are relatively well understood, comparatively little is known about H_2 +CH₄ mixed hydrates.

 H_2 was initially believed to be too small to stabilize hydrate cavities, and it was not until 1999 that the phase diagram of the H_2 hydrate system was reported by Dyadin *et al.* [6]. Since this

pioneering work, several other key discoveries have been made highlighting the structure, cage occupancies, and storage capacities of H₂ hydrates [7–24]. Despite encouraging initial results, the major consensus of much of the previous work has been that due to the extremely high pressures and/or low temperatures (e.g. $p \approx 150$ MPa, T = 270 K) required for their stabilization [11], simple H₂ hydrates may be impractical for application.

As a means of easing the extreme thermodynamic requirements, several studies have investigated the possibility of adding small amounts of a second larger guest to better stabilize the hydrate structure [9,10,12–15,17,18,20,21,23,24]. One of the more encouraging binary hydrate systems is H₂+THF mixed hydrate. By simply adding 5.6 mol% THF, mixed hydrate can form at much more moderate conditions than simple H₂ hydrate [13]. Specifically, simple H₂ hydrates form at around p = 150 MPa and T = 270 K [11], while H₂+THF mixed hydrates form at around p = 10MPa and T = 280 K [9,13]. However, the trade off in this scenario is that only a limited amount of H₂ can be enclathrated in the mixed hydrate as stated in Chapter II Section 1. For hydrates to become a practical storage medium, it is important to achieve a high energy density at relatively mild conditions. Addressing this issue, Lee et al. and Sugahara et al. reported a "tuning effect" for the H₂+THF hydrate system in 2005 and 2009, respectively [10,21]. They reported that by reducing the amount of THF added to below 5.6 mol%, H₂ could occupy not only the S-cages but also some of the L-cages. In principle, tuned H₂+THF hydrates could be an interesting energy storage medium. The proof-of-concept for the tuned H₂+THF hydrates was confirmed in Chapter II Section 1. However, reproducing the quantitative results on the tuning effect for the H₂+THF system has been a source of controversy [13,14,20,23].

I suggest that, like THF, CH₄ may also serve as a thermodynamic promoter, but unlike THF, CH₄ is small enough that H₂ may compete for occupancy in the larger cages resulting in an overall increase in H₂ enclathration. Furthermore, H₂+CH₄ mixed hydrates are expected to have the higher overall energy density than H₂+THF hydrates because CH₄ serves as an additional source of energy, unlike THF. When compared to H₂+THF mixed hydrate, hydrates incorporating H₂ and CH₄ are comparatively underexplored [15,22,24], with some authors suggesting the H₂+CH₄ system as incapable of forming binary hydrates [15]. In this section, I confirm the formation of binary H₂+CH₄ hydrates and report preliminary results on the thermodynamics of this system, including the effect of initial mole fraction of mixed gas, pressure, and formation period. Hydrate crystal structure and cage occupancies of guest species were obtained from PXRD and Raman spectroscopy, respectively.

II.2.3. Experimental

Apparatus

A high-pressure vessel (Hip, inner volume: $\sim 4 \text{ mL}$) was used for hydrate preparation. A gas chromatograph (GC) (Hewlett Packard, 5890 Series II) with a thermal conductivity detector (TCD-GC) was used for decision of gas composition. A powder X-ray diffractometer (Siemens, Kristalloflex 805) with Cu X-ray (generation power: 45 kV, 40 mA) was used for determination of crystal structure of hydrate. A Raman spectrometer (Horiba, Jobin Yvon) with a diode laser (wavelength: 532.268 nm) was used for clarification of cage occupancy of guest species.

Procedure

Hydrates studied in this study were synthesized by pressurizing ice powders incorporating particles less than 180 µm. About 1 g of the ice powder was loaded into a high-pressure cell of internal volume ~ 4 mL followed by immersion in a glycol cooling bath held at 263 K. The cell containing the ice powder was allowed to equilibrate in the bath for more than an hour so that the inner cell temperature would rise from LN_2 temperature to the bath temperature. The cell was then pressurized first with CH₄ followed by H₂ up to the respective desired pressures for each experiment. The partial pressures of each component were calculated using the Peng-Robinson equation of state [25,26] to satisfy the desired composition of the initial mixed gas. As a means of confirming the initial gas composition, an empty cell was simultaneously pressurized in parallel with the same mixed gas and the contents in the empty cell were later analyzed with GC in order to determine any differences between the measured and estimated mole fractions. After pressurization, the cell was left in the glycol bath for a desired formation period. Then the cell was quenched in LN₂ for 15 minutes and vented. The hydrate sample was then analyzed with PXRD and Raman spectrometer at p = 0.1 MPa and T = 83 K. PXRD measurement was performed in the step-scan mode with a dwell time of 2 s and a step size of 0.02°. The PXRD pattern indexing and cell refinement were performed with PowderX [27] and Chekcell [28] programs.

II.2.4. Results and Discussion

Whereas the thermodynamics of binary hydrates including H₂ and larger guest molecules such as THF has been well characterized [9,10,12–14,17,18,20,21,23], comparatively little is known about the thermodynamics of H₂+CH₄ binary hydrate [15,22,24]. From the available studies, the general agreement is that at modest mole fractions of CH₄ [22], s-I is the preferred structure [15,24]. However, it is expected that as the mole fraction of CH_4 decreases, s-II may become more favorable as simple H_2 hydrates naturally form s-II [7]. In the following figures, I show that like other known gas mixtures such as $CH_4+C_2H_6$ [29–31], the hydrate structure of the H_2+CH_4 mixed system is dependent on at least three variables; initial gas mole fraction, pressure and formation period.

Figures II-2-1 and 2 show Raman spectroscopic evidence of hydrate structural dependence on the initial composition of the gas mixture and total system pressure when held at constant temperature (T = 263 K) and formation period (t = 72 h). At a constant pressure (p = 70 MPa), samples prepared at $y_{CH4} \le 0.01$ formed no hydrate, $0.03 \le y_{CH4} \le 0.05$ formed s-II hydrates, and $y_{CH4} \ge 0.07$ formed s-I hydrates. At a fixed concentration ($y_{CH4} = 0.05$), the results indicate that s-I hydrates are initially stable starting around p = 50 MPa and it is only when the pressure exceeds $p \approx$ 70 MPa does s-II become the more favorable structure.



Figure II-2-1. Raman spectra indicating composition dependence at p = 70 MPa and t = 72 h.



Figure II-2-2. Raman spectra indicating initial pressure dependence at $y_{CH4} = 0.05$ and t = 72 h.

Hydrate structure was confirmed directly by PXRD as well as indirectly by the Raman shift. In **Figure II-2-1** at $y_{CH4} \ge 0.07$, the C–H stretching vibration mode region shows two peaks occurring at $\Delta \nu = 2901$ cm⁻¹ and 2913 cm⁻¹ corresponding with CH₄ occupancy of the M- and Scages, respectively, in s-I hydrates [24]. Furthermore, the approximate peak area ratio (A_L/A_S) of the two contributions is 3.26, which is similar to the stoichiometry of the s-I hydrate unit cell (two Scages + six M-cages). Conversely, at $y_{CH4} \le 0.05$, the approximate peak area ratio (A_L/A_S) of 0.55 confirms the hydrate structure had changed to s-II (sixteen S-cages + eight L-cages).

In contrast to the C–H stretching vibration mode region, where only two peaks are observed (one per cage), in the H–H stretching vibration mode region, multiple peaks per cage are observed due to the multiple clusters and quantum spin of H₂ [19]. H₂ has two kinds of spin isomers, ortho and para [32], occurring in a ratio 3:1 at temperatures near the ice point. For s-I hydrates, the peaks at $\Delta \nu = 4121$ cm⁻¹ and 4127 cm⁻¹ are derived from H₂ singly occupied in S-cages, the peaks at $\Delta \nu = 4126$ cm⁻¹ and 4132 cm⁻¹ are from H₂ singly occupied in M-cages, and the peaks at $\Delta \nu = 4144$ cm⁻¹ and 4150 cm⁻¹ indicate H₂ doubly occupied in M-cages, as previously reported [24]. In the case of s-II hydrate, the peaks at $\Delta \nu = 4121$ cm⁻¹ and 4136 cm⁻¹ are derived from doubly occupied L-cages, the peaks at $\Delta \nu = 4138$ cm⁻¹ and 4144 cm⁻¹ indicate triply occupied L-cages, and the peaks at $\Delta \nu = 4146$ cm⁻¹ and 4152 cm⁻¹

From the PXRD measurements in **Figure II-2-3**, the space group and lattice constant of a typical s-I H₂+CH₄ hydrate sample were determined to be *Pm3n* with $a = 1.187 \pm 0.001$ nm, and for a typical s-II H₂+CH₄ hydrate, *Fd3m* and $a = 1.715 \pm 0.003$ nm. Interestingly, some of the s-I and s-II hydrate samples showed a significant amount of ice Ih in coexistence with the hydrate phase despite initially being well inside the estimated s-II hydrate phase space (**Figure II-2-4**). This appearance of ice is likely a result of isochoric procedure used in synthesizing the hydrates. Specifically, although the initial gas phase composition is within the thermodynamically stable region for s-II hydrate, due to the larger molecular diameter of CH₄, CH₄ may preferentially become enclathrated, resulting in a depletion of available CH₄ in the gas phase moving the system towards the equilibrium conditions (H_{II}S₁G) after a small fraction of conversion.



Figure II-2-3. PXRD pattern of three hydrate samples obtained in the present study highlighting the characteristic peaks for s-I, s-II, and hydrate free samples (ice-Ih).



Figure II-2-4. Experimentally observed hydrate structures over a range of pressures and initial gas compositions in H_2 +CH₄+water system at t = 72 h and T = 263 K.

From the viewpoint of H_2 storage capacity, s-II hydrate is expected to be more desirable than s-I hydrate due to the ability of the L-cage of s-II hydrate to enclathrate up to four H_2 molecules compared to two H_2 in the M-cage of s-I hydrate [16]. However, from **Figures II-2-1 and 2**, s-II hydrates were observed only in a limited range of conditions. Consequently, a series of additional experiments examining the hydrate structure at a wider range of pressures and CH₄ concentrations were performed to provide a better estimate of the phase space, as shown in **Figure II-2-4**.

Figure II-2-4 shows that at T = 263 K four distinct regions are present in the H₂+CH₄ system in the range of p = 10–80 MPa and $y_{CH4} = 0$ –0.12. Some samples, for example measured at $y_{CH4} =$ 0.10 and p = 70 MPa, showed the co-existence of s-I and s-II hydrates, this implying the system was not yet at equilibrium and maybe a result of the isochoric method. These initial results reveal that from a practical standpoint, although s-II hydrate may have a higher H₂ density, the thermodynamic window for s-II hydrate is small and the requirements necessary for formation, $p \ge$ 70 MPa, remain extreme even after the addition of a CH₄ co-guest molecule. Furthermore, as the pressure decreases around p = 20 MPa, the driving force required to enclathrate H₂ in either structure was too low, resulting in the formation of simple CH₄ hydrate. In one additional study, the effect of formation period at a fixed mole fraction ($y_{CH4} = 0.03$) and pressure (p = 70 MPa) was studied (**Figure II-2-5**). Interestingly, although s-II was determined to be the thermodynamically preferred structure at these conditions (as shown above in **Figures II-2-1 and 4**), a metastable s-I hydrate phase was observed as the kinetically favorable structure in the early stages of formation. It was only at t = 5 h that the spectra indicate a change from s-I hydrate to s-II hydrate. To explain this structural change during the formation process, I propose that the ice initially surrounded with CH₄ will quickly convert to a metastable s-I CH₄-rich hydrate seed. Then, the H₂+CH₄ hydrate may then convert to the equilibrium s-II structure with the spectra appearing mostly constant after t = 24 h.



Figure II-2-5. Raman spectral change for each formation period at $y_{CH4} = 0.03$ and p = 70 MPa.

II.2.5. Conclusions

H₂+CH₄ hydrates prepared at various conditions were analyzed by PXRD and Raman spectroscopy to reveal hydrate structure and cage occupancies of H₂ and CH₄, respectively. The structure of H₂+CH₄ hydrate was determined to be strongly dependent on the mole fraction of mixed gas, total system pressure, and formation period. H₂ and CH₄ were observed to competitively occupy the S-, M-, and L-cages. These hydrates form at more moderate conditions compared to simple H₂ hydrate and H₂+THF tuned hydrate.

[-]

II.2.6. Nomenclatures

θ	Diffraction angle [°]
ν	Wavenumber [m ⁻¹]
$A_{ m L}$	Raman peak area corresponding CH4 occupying large cage [-]
As	Raman peak area corresponding CH4 occupying small cage [-]
a	Lattice constant [m]
р	Pressure [Pa]
Т	Temperature [K]
t	Formation period [h]
УСН4	Mole fraction of CH ₄ in gas phase (water free) [-]
ZCH4	Mole fraction of CH ₄ in hydrate phase (water free) [-]
G	Gas phase
H _{II}	s-II hydrate phase
S_1	Ice phase

II.2.7. References

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Section 3

Isothermal Phase Equilibria for the Difluoromethane+1,1,1,2-Tetrafluoroethane Mixed-Gas Hydrate System

II.3.1. Abstract

Isothermal phase equilibria (pressure-composition relations in HL₁G) in the difluoromethane (HFC-32)+1,1,1,2-tetrafluoroethane (HFC-134a) mixed-gas hydrate system were measured at T = 274.15 K, 279.15 K, and 283.15 K. The heterogeneous azeotropic-like behavior derived from the structural phase transition of HFC-32+HFC-134a mixed-gas hydrates appears over the whole temperature range of the present study. In addition to the heterogeneous azeotropic-like behavior, the isothermal phase equilibrium curves of the HFC-32+HFC-134a mixed-gas hydrate system exhibit the negative homogeneous azeotropic-like behavior at T = 279.15 K and 283.15 K. The negative azeotropic-like behavior, which becomes more remarkable at higher temperatures, results in the lower equilibrium pressure of HFC-32+HFC-134a mixed-gas hydrate sthan that of both simple HFC-32 and HFC-134a hydrates. Although the simple HFC-134a hydrate is transformed from s-II hydrate to s-I hydrate at p = 65 MPa and T = 282.2 K, the present findings reveal that HFC-134a molecules occupy a part of the M-cages of the s-I mixed-gas hydrate even at pressures much lower than p = 65 MPa.

II.3.2. Introduction

Hydrates get much attention as newly refrigerant materials because gas hydrates have appropriate properties for cold energy storage [1,2]. HFCs are well-known refrigerant materials due to the thermodynamic characteristics and chemical stabilities. The hydrates prepared from HFCs have higher enthalpy than liquefied HFCs [3,4]. The phase equilibrium relations and the dissociation enthalpies for some simple HFC hydrates have been reported [5–7]. The studies on HFCs mixed-gas hydrate systems, however, are rarely reported.

In this section, isothermal phase equilibria (pressure-composition) in an HFCs mixed-gas hydrate system including typical refrigerant gases, HFC-32 and HFC-134a, were investigated. The main purpose of the present study is to reveal the three-phase equilibrium relations of HL₁G in the HFC-32+HFC-134a mixed-gas hydrate system for the improvement of refrigerant cycles. In addition, the hydrate structures and the cage occupancies of guest species are discussed, based on the Raman spectra.

II.3.3. Experimental

Materials

The materials used in the present study are summarized in **Table II-3-1**. All of them were used without further purification.

Material nameSourceMole fraction purityHFC-32Neriki Gas Co., Ltd.> 0.999HFC-134aDaikin Ind., Ltd.> 0.996Distilled waterWako Pure Chemicals Ind., Ltd.> 0.9999

Table II-3-1. Information on the materials used in the present study.

Apparatus

The present experimental apparatus for the isothermal phase equilibrium measurement is shown in **Figure II-3-1**. A high-pressure cell (inner volume: 150 cm³, maximum working pressure: 10 MPa) was used. The temperature was controlled with a thermocontroller (Taitec, CL-80R). The equilibrium pressure was measured with a pressure gauge (Valcom, VPRT, maximum uncertainty: 0.004 MPa). The equilibrium temperature was measured with a thermistor thermometer (Takara, D632, reproducibility: 0.02 K). A TCD-GC (Shimadzu, GC-14B, maximum uncertainty: 0.005) was used for determination of compositions. The pressure was calibrated by a Ruska quartz Bourdon tube gauge (Ruska, Pressure Counter 3855-703-00). The temperature was calibrated with a platinum resistance thermometer (25 Ω) defined by ITS-90.



Figure II-3-1. Schematic illustration of the experimental apparatus for phase equilibrium measurement performed in the present study.

The present experimental apparatus for the Raman spectroscopic analysis is shown in **Figure II-3-2**. The high-pressure optical cell (inner volume: 0.1 cm^3 , maximum working pressure: 400 MPa) was used for Raman spectroscopy. A programming thermocontroller (Taitec, CL-80R) adjusted the cell temperature. The equilibrium pressure was measured by a pressure gauge (Valcom, VPRT, maximum uncertainty: 0.02 MPa). The equilibrium temperature was measured with the thermistor thermometer (Takara, D641, reproducibility: 0.02 K). The pressure and temperature were calibrated with as the same ways as mentioned above. A laser Raman microprobe spectrometer with a multichannel CCD detector (Horiba, Jobin-Yvon T64000) and an argon ion laser beam (514.5 nm, 100 mW) condensed to 2 μ m in spot diameter were used to analyze hydrate single crystals. The CCD detector was maintained at *T* = 140 K by LN₂ for heat-noise reduction. The spectral resolution was approximately 0.7 cm⁻¹. The Raman shift was calibrated with the neon emission lines in the air.



Figure II-3-2. Schematic illustration of the experimental apparatus for Raman spectroscopy performed in the present study.

Procedure

For phase equilibrium measurements, the gas mixture was introduced into an evacuated highpressure cell. Distilled water was successively introduced into the cell to pressurize up to a desired pressure. The cell was immersed in a thermostatic bath for controlling the system temperature precisely. The content was cooled and agitated by a mixing bar that is moved up-and-down by magnetic attraction from outside of the cell. The phase behavior was directly observed through sapphire windows attached to the cell. After the formation of the mixed-gas hydrates, the temperature was kept constant for at least one day to establish the three-phase equilibrium conditions. After that, a small amount of samples separately taken from the gas and aqueous phases was analyzed by means of GC. The experimental procedure was repeated with various compositions to obtain the three-phase equilibrium (pressure-composition) relations. The equilibrium compositions of hydrate phase were measured in the following way. The mixed-gas hydrates under three-phase equilibrium conditions were prepared with the same procedure mentioned above. Then, only the excess water was sufficiently drained, and then the mixed-gas hydrates were quenched for more than 6 h at T = 263 K. The quenched mixed-gas hydrates were replaced from the formation cell to another one in the refrigerated room controlled at T = 263 K. The sample was allowed to dissociate under room temperature and atmospheric pressure conditions. The released mixed gas was analyzed by means of GC.

For Raman spectroscopy, the mixed-gas hydrates were generated in a high-pressure optical cell by the same procedure mentioned in the previous paragraph. The content was cooled and agitated by an enclosed ruby ball. The temperature was controlled by circulating thermostatic water through the jacket of the cell. After the formation of the mixed-gas hydrates, the temperature was gradually increased and decreased to prepare hydrate single crystals under three-phase coexisting conditions. Then the single crystals were annealed to avoid metastability, and the temperature was kept constant for more than one day to establish the three-phase equilibrium conditions. The phase behavior was observed with CCD camera through the sapphire window. The hydrate single crystal was analyzed by use of Raman spectrometer with multichannel CCD detector. The argon ion laser beam was irradiated from the object lens to the hydrate single crystal through the upper sapphire window.

II.3.4. Results and Discussion

Isothermal pressure (p)-composition (x_{HFC-32} , y_{HFC-32} , z_{HFC-32}) diagrams for the HFC-32+HFC-134a mixed-gas hydrate system at T = 274.15 K, 279.15 K, and 283.15 K under three-phase equilibrium conditions of hydrate, gas, and aqueous phases are shown in Figures II-3-3 to 5, respectively. The horizontal axis in Figures II-3-3 to 5 indicates the equilibrium composition of HFC-32 on a water-free basis. Table II-3-2 summarizes the $p-x_{HFC-32}$, y_{HFC-32} relations for each isotherm obtained without the dissociation of hydrate samples. The obviously unreasonable data in the aqueous phase were removed, due to the incidental involvement of hydrate crystal in the sample of aqueous phase. Table II-3-3 summarizes the $p-z_{\rm HFC-32}$ relation for each isotherm. As shown in Figures II-3-4 and 5, the isothermal phase equilibrium relations of the HFC-32+HFC-134a mixedgas hydrate system exhibit the negative homogeneous azeotropic-like behavior at T = 279.15 K and 283.15 K. The local pressure minima are located at $p = 0.159 \pm 0.004$ MPa, $y_{HFC-32} = 0.175 \pm 0.005$ at $T = 279.15 \pm 0.02$ K and $p = 0.302 \pm 0.004$ MPa, $y_{HFC-32} = 0.471 \pm 0.005$ at $T = 283.15 \pm 0.02$ K. Especially in the case of T = 283.15 K, a large pressure reduction from the equilibrium pressures of both the simple HFC-32 and HFC-134a hydrates is useful and effective for the refrigerant systems. The pressure reduction is getting small with the decrease in temperature, and finally, the azeotropiclike behavior disappears at T = 274.15 K. Also, the local pressure minimum moves to lower composition range with the decrease in temperature. The experimentally-observed negative azeotropic-like behavior has been already reported in the some mixed-gas hydrate systems, for example, s-II CH₄+C₃H₈ [8], s-II krypton (Kr)+C₃H₈ [8], and s-I CH₄+C₂H₄ [9] mixed-gas hydrate systems. The s-I CH₄+C₂H₄ mixed-gas hydrate system exhibits negative azeotropic behavior at T =

293 K, while the azeotropic behavior disappears at T = 284 K [9]. These results suggest that, at a relatively high temperature, the azeotropic-like behavior is clearly observed. Note that the HFC-32+HFC-134a binary system (excluding water) exhibits no azeotropic-like behavior on the vapor-liquid equilibrium relation in a temperature range of the present study [10,11].



Figure II-3-3. Isothermal phase equilibria of hydrate, gas, and aqueous phases in the HFC-32+HFC-134a mixed-gas hydrate system at T = 274.15 K.



Figure II-3-4. Isothermal phase equilibria of hydrate, gas, and aqueous phases in the

HFC-32+HFC-134a mixed-gas hydrate system at T = 279.15 K.



*X*_{HFC-32}, *Y*_{HFC-32}, *Z*_{HFC-32}

Figure II-3-5. Isothermal phase equilibria of hydrate, gas, and aqueous phases in the HFC-32+HFC-134a mixed-gas hydrate system at T = 283.15 K.

<i>p</i> / MPa ^{<i>a</i>}	YHFC-32 ^a	XHFC-32 ^a	<i>p</i> / MPa ^a	YHFC-32 ^a	XHFC-32 ^a	<i>p</i> / MPa ^a	YHFC-32 ^a	X HFC-32 a
$T = 274.15 \text{ K}^{a}$			$T = 279.15 \text{ K}^{a}$			$T = 283.15 \text{ K}^{a}$		
0.056	0.000	0.000	0.167	0.000	0.000	0.412	0.000	0.000
0.060	0.167	0.286	0.164	0.025	0.049	0.396	0.041	0.160
0.065	0.293	0.482	0.159	0.224	0.394	0.383	0.060	0.347
0.070	0.410	0.608	0.161	0.358	0.592	0.344	0.150	0.433
0.081	0.637	0.790	0.165	0.456	0.692	0.319	0.254	0.642
0.105	0.803	0.883	0.173	0.558	0.788	0.311	0.319	0.689
0.134	0.870	0.936	0.196	0.706	0.872	0.303	0.415	0.765
0.147	0.903	0.955	0.224	0.803	0.931	0.302	0.471	_b
0.171	0.930	0.983	0.259	0.870	0.944	0.307	0.553	_b
0.177	0.938	0.988	0.302	0.918	0.976	0.345	0.745	0.895
0.175	0.946	_b	0.313	0.928	0.982	0.381	0.826	0.974
0.170	0.979	0.995	0.307	0.954	0.988	0.422	0.870	0.959
0.169	1.000	1.000	0.293	1.000	1.000	0.491	0.915	_b
						0.486	0.928	0.983
						0.461	1.000	1.000

Table II-3-2. Isothermal phase equilibrium data of gas and aqueous phases coexisted with HFC-32+HFC-134a mixed-gas hydrates.

^{*a*} u(p) = 0.004 MPa, u(T) = 0.02 K, and $u(y_{HFC-32}) = u(x_{HFC-32}) = 0.005$.

^{*b*} The only *p*-*y*_{HFC-32} relation is listed in spite of the lack of $x_{\text{HFC-32}}$ because the datum set is still meaningful for understanding the phase behavior.

<i>p</i> / MPa ^{<i>a</i>}	ZHFC-32 ^{<i>a</i>}	<i>p</i> / MPa ^{<i>a</i>}	ZHFC-32 ^a	<i>p</i> / MPa ^{<i>a</i>}	ZHFC-32 ^a
$T = 274.15 \text{ K}^{a}$		T = 279	$T = 279.15 \text{ K}^a$.15 K ^a
0.06	0.00	0.17	0.00	0.41	0.00
0.07	0.27	0.16	0.08	0.35	0.20
0.10	0.55	0.16	0.20	0.33	0.29
0.12	0.62	0.17	0.48	0.33	0.57
0.16	0.70	0.19	0.60	0.37	0.68
0.17	1.00	0.20	0.65	0.40	0.74
		0.26	0.71	0.48	0.81
		0.30	0.76	0.48	0.99
		0.29	1.00	0.46	1.00

Table II-3-3. Isothermal pressure-composition relations in hydrate phase for theHFC-32+HFC-134a mixed-gas hydrate system.

^{*a*} u(p) = 0.004 MPa, u(T) = 0.02 K, and $u(z_{HFC-32}) = 0.01$.

Holder and Manganiello [12] and Thakore and Holder [8] claimed that the larger stability of hydrates with certain compositions is due to competing effect of small and large guest molecules and the hydrate structure (they claimed that no azeotropic behavior appears in s-I hydrates [8]), not a non-ideality of hydrates. The results in the present study indicate that the occurrence of the azeotropic behavior cannot be explained only by competing effect due to different sizes of both guest species. The temperature effect on the cage occupancies of both guest species has to be considered.

In addition to the homogeneous azeotropic-like behavior, the heterogeneous azeotropic-like behavior appears in the HFC-32+HFC-134a mixed-gas hydrate system. As shown in **Figures II-3-3** to **5**, a quadruple point exists at $y_{HFC-32} \approx 0.9$ in the equilibrium p- y_{HFC-32} curve and the pressures are higher than equilibrium ones of both simple HFC-32 and HFC-134a hydrates at the same temperature. In general, the structural phase transition of hydrates properly occurs in the mixed-gas hydrate when the crystal structures of the simple hydrates are different from each other. To demonstrate that the heterogeneous azeotropic-like behavior is derived from the structural phase transition, the Raman spectra corresponding to the intramolecular vibration modes of the HFC-32 and HFC-134a molecules at T = 283.15 K are shown in **Figure II-3-6**. The top spectrum at $y_{HFC-32} = 0.98$ is completely different from others and very similar to that of s-I simple HFC-32 hydrate [4],

while the spectra at $y_{HFC-32} = 0.26$, 0.80, and 0.89 are similar to the bottom spectrum derived from the s-II simple HFC-134a hydrate. The Raman spectra reveal the structural phase transition occurs at the quadruple point (H_IH_{II}L₁G). The details of the quadruple points (H_IH_{II}L₁G) are summarized in **Table II-3-4**. The quadruple point (H_IH_{II}L₁G) is one of the most characteristics in the present mixed-gas hydrate system.



Figure II-3-6. Raman spectra of HFC-32 and HFC-134a molecules in the HFC-32+HFC-134a mixed-gas hydrates at T = 283.15 K, and Raman spectrum in the simple HFC-134a hydrate at T = 279.15 K and p = 0.17 MPa. The Raman peaks detected around $\Delta \nu = 840$ cm⁻¹, 1090 cm⁻¹, and 2975 cm⁻¹ correspond to the intramolecular vibration modes of HFC-134a molecule (broken lines). The peaks detected around $\Delta \nu = 1098$ cm⁻¹, 2835 cm⁻¹, 2950 cm⁻¹, and 3020 cm⁻¹ are the intramolecular vibration modes of HFC-32 molecule in s-I hydrates divide into doublet except for that of $\Delta \nu = 1098$ cm⁻¹ (C–F stretching vibration mode) [4].

T / K a	<i>p</i> / MPa ^{<i>a</i>}	УНFС-32 ^а	X HFC-32 a	$Z_{\rm HFC-32}({\rm s-II})^a$
274.15	0.177	0.938	0.988	0.73
279.15	0.313	0.928	0.982	0.77
283.15	0.491	0.915	0.982	0.83

Table II-3-4. Summary on the structural phase transition points in the HFC-32+HFC-134a mixedgas hydrate system. The value of $z_{\text{HFC-32}}$ in the s-I hydrate phase is speculated to be more than 0.99 on each isotherm.

 $^{a}u(p) = 0.004$ MPa, u(T) = 0.02 K, $u(y_{HFC-32}) = u(x_{HFC-32}) = 0.005$, and $u(z_{HFC-32}) = 0.01$.

The Raman spectra provide the additional knowledge regarding the cage occupancy of guest species. HFC-134a molecules occupy only L-cages of the s-II HFC-32+HFC-134a mixed-gas hydrates in the composition range of $y_{HFC-32} = 0.26-0.89$, based on the single Raman peak in the simple HFC-134a hydrate shown in bottom spectrum of Figure II-3-6. In the hydrate equilibrated with the gas phase of $y_{HFC-32} = 0.98$, a weak Raman peak was detected at $\Delta v = 847$ cm⁻¹. This Raman shift does not correspond to the C-C stretching vibration mode of HFC-134a enclathrated into L-cages ($\Delta v = 841 \text{ cm}^{-1}$) of s-II hydrates. It is also different from the Raman shift of HFC-134a involved in gas phase ($\Delta v = 843$ cm⁻¹) as shown in Figure II-3-7. At the present experimental temperatures and pressures, HFC-134a molecules occupy only the L-cages of s-II simple HFC-134a hydrate. As a supplemental experiment, phase equilibrium curve of HL₁G for the simple HFC-134a hydrate system and cage occupancy of HFC-134a were investigated by means of phase equilibrium measurement and Raman spectroscopy, respectively. Phase equilibrium relation of simple HFC-134a hydrate system is shown in Figure II-3-8 and summarized in Table II-3-5. Raman spectrum obtained from the simple HFC-134a hydrate is also shown in Figure II-3-7. The result is that the structural phase transition point between s-I and s-II hydrates exists at $p = 65 \pm 1$ MPa, T = 282.2 ± 0.1 K and HFC-134a occupies M-cages of s-I hydrate above the structural phase transition point. Therefore, Raman peak detected at $\Delta v = 847$ cm⁻¹ reveals that the coexistence of HFC-32 molecule in the hydrate crystal results in the M-cage occupancy of HFC-134a molecule. The Mcage occupancy of the HFC-134a molecule leads to the heterogeneous azeotropic-like behavior, that is, the pressure-increase from the equilibrium pressure of the simple HFC-32 hydrate.



Figure II-3-7. Raman spectra of the C–C stretching vibration mode of HFC-134a molecule between hydrate at $y_{HFC-32} = 0.89$ and 0.98 and gas phases.



Figure II-3-8. Three-phase equilibrium relations for the HFC-134a+water binary system.

HFC-32 molecule is enclathrated in S- and M-cages of the s-I simple HFC-32 hydrate [4]. Raman spectrum in the s-I HFC-32+HFC-134a mixed-gas hydrate at $y_{\text{HFC-32}} = 0.98$ is similar to that of the simple HFC-32 hydrate [4]. At $y_{\text{HFC-32}} = 0.26$ to 0.89, Raman peaks of the HFC-32 molecules in the L-cages of the s-II HFC-32+HFC-134a mixed-gas hydrates do not seem to be detected. However, a small amount of HFC-32 molecules should occupy the L-cages of the s-II HFC-32+HFC-134a mixed-gas hydrates in the certain composition range because the mole fraction of HFC-32 in hydrate phases is beyond 2/3 (the ratio $N_{\text{S}}/(N_{\text{S}}+N_{\text{L}})$ in the unit cell of s-II hydrates).

<i>T /</i> K	<i>p</i> / MPa
283.12	10.63
283.14	18.32
283.04	20.02
282.93	30.03
282.82	34.75
282.77	42.04
282.63	49.73
282.53	54.77
282.27	65
282.79	70
283.97	81
284.70	87
286.37	100
288.02	116
290.28	141
291.69	156
292.78	171
294.07	187
297.51	229
298.69	245
299.96	266
301.62	293

Table II-3-5. Three-phase (HL₁L₂) equilibrium data for the HFC-134a+water system.^{*a*} The structural phase transition point is located at $p = 65 \pm 1$ MPa and $T = 282.2 \pm 0.1$ K.

 $^{a}u(p) = 0.15$ MPa at p < 60 MPa, u(p) = 2 MPa at p > 60 MPa, and u(T) = 0.02 K.

II.3.5. Conclusions

The isothermal phase equilibrium relations of the HFC-32+HFC-134a mixed-gas hydrate system were investigated under the three-phase equilibrium conditions. The remarkable characteristics in the HFC-32+HFC-134a mixed-gas hydrate system are that the isothermal phase equilibrium curves exhibit both the homogeneous and heterogeneous azeotropic-like behavior at T = 279.15 K and 283.15 K, while they exhibit only the heterogeneous azeotropic-like behavior at T = 274.15 K. This negative homogeneous azeotropic-like behavior enables the preparation of HFC-32+HFC-134a mixed-gas hydrates under low-pressure conditions. In addition, Raman spectra reveal that the HFC-134a molecule, in spite of an s-II hydrate former, occupies the M-cage of the s-I mixed-gas hydrate.

II.3.6. Nomenclatures

ν	Wavenumber [m ⁻¹]
$N_{ m L}$	Number of L-cages in the unit cell of s-II hydrates [-]
Ns	Number of S-cages in the unit cell of s-II hydrates [-]
р	Pressure [Pa]
Т	Temperature [K]
и	Standard uncertainties of pressure [Pa], temperature [K], and mole fraction [-]
XHFC-32	Mole fraction of HFC-32 in aqueous phase (water free) [-]
YHFC-32	Mole fraction of HFC-32 in gas phase (water free) [-]
ZHFC-32	Mole fraction of HFC-32 in hydrate phase (water free) [-]
G	Gas phase
Н	Hydrate phase
H_{I}	s-I hydrate phase
H_{II}	s-II hydrate phase
L_1	Aqueous phase
L_2	HFC-134a-rich liquid phase

II.3.7. References

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Section 4

Isothermal Phase Equilibria for the Carbon Dioxide+1,1,1,2-Tetrafluoroethane and Carbon Dioxide+1,1-Difluoroethane Mixed-Gas Hydrate Systems

II.4.1. Abstract

Isothermal phase equilibrium relations of the CO₂+HFC-134a and CO₂+1,1-difluoroethane (HFC-152a) mixed-gas hydrate systems were measured at T = 279.15 K. The isothermal phase equilibria of the CO₂+HFC-134a mixed-gas hydrate system exhibit the heterogeneous azeotropic-like behavior derived from the structural phase transition of the mixed-gas hydrate. The CO₂+HFC-152a mixed-gas hydrate system shows neither a structural phase transition nor azeotropic-like behavior.

II.4.2. Introduction

Unlike simple gas hydrate systems, the particular mixed-gas hydrate system has characteristic phenomena that depend on the composition of guest mixtures, that is, the structural phase transition and the azeotropic-like behavior. In general, the structural phase transition of hydrates surely occurs in the mixed-gas hydrate when the structures of the simple gas hydrates are different from each other. In addition to the commonsensical transition, another structural phase transition has been reported, which the s-II mixed-gas hydrates emerge in a certain composition region; nevertheless each guest molecule generates an s-I hydrate. The latter type of structural phase transition has been reported only in the $CO_2+c-C_3H_6$ mixed-gas hydrate system [1] among the CO_2 -containing mixed-gas hydrate systems. It does not occur in the hydrate systems of the $CO_2+C_2H_6$ and $CO_2+trifluoromethane (CHF_3)$ [2].

The azeotropic-like behavior has been experimentally observed in some mixed-gas hydrate systems. For example, in the HFC-32+HFC-134a mixed-gas hydrate system as shown in **Chapter**

II Section 3, both the negative homogeneous azeotropic-like behavior and the heterogeneous azeotropic-like behavior is observed at T = 279.15 K and 283.15 K.

HFCs such as HFC-152a and HFC-134a are still key components as countermeasures against destruction of the ozone layer, although it is a temporary expedient until some reliable substitute is found in the associated industries. The mixed-gas hydrate involving HFCs is a favorable heat-conducting medium in the air-conditioning system for reducing the amount of HFCs. In addition, if the system exhibits the negative azeotropic-like behavior, the operational conditions would become mild because the minimum equilibrium pressure is depressed from those of both simple gas hydrates.

HFC-152a is as large as c-C₃H₆, which is one of the largest guest species forming s-I hydrates [3]. As described above, the structural phase transition occurs in the CO₂+c-C₃H₆ mixed-gas hydrate system [1]. If the size of guest species were the most important for the structural phase transition, a possible structural phase transition in the CO₂+HFC-152a mixed-gas hydrate system would be implied.

In this section, to clarify whether the CO₂+HFC-152a and CO₂+HFC-134a mixed-gas hydrate systems exhibit such characteristic phenomena, I have investigated isothermal phase equilibria for the mixed-gas hydrates under the three-phase (HL₁G) equilibrium conditions at T = 279.15 K. In addition, the single crystal of mixed-gas hydrates has been analyzed by *in situ* Raman spectroscopy.

II.4.3. Experimental

Materials

The materials used in the present study are summarized in **Table II-4-1**. All of them were used without further purification.

Material name	Source	Mole fraction purity
CO ₂	Neriki Gas Co., Ltd.	> 0.9999
HFC-152a	Daikin Ind., Ltd.	> 0.995
HFC-134a	Daikin Ind., Ltd.	> 0.996
Distilled water	Wako Pure Chemicals Ind., Ltd.	> 0.9999

Table II-4-1. Information on the materials used in the present study.

Apparatus

The experimental apparatus for isothermal phase equilibrium measurement and Raman spectroscopy used in the present study are the same as described in **Chapter II Section 3**.

Procedure

The experimental procedures for isothermal phase equilibrium measurement and Raman spectroscopy performed in the present study are the same as described in **Chapter II Section 3**. As a typical example, a photo of the CO₂+HFC-152a mixed-gas hydrate crystals in the high-pressure cell at p = 0.23 MPa and T = 279.15 K is shown in **Figure II-4-1**.



Figure II-4-1. CO₂+HFC-152a mixed-gas hydrate crystals generated under the three-phase coexisting conditions at p = 0.23 MPa and T = 279.15 K. (Many crystals are observed in the aqueous solution. A spherical ruby is trapped in the high-pressure cell for stirring the gas-liquid interface.)

II.4.4. Results and Discussion

CO₂+HFC-134a Mixed-Gas Hydrate System

Isothermal phase equilibrium relations for the CO₂+HFC-134a mixed-gas hydrate system at T = 279.15 K are shown in **Figure II-4-2** and summarized in **Table II-4-2**. The equilibrium CO₂ compositions in the gas and aqueous phases at a given pressure are comparable and larger than that of hydrate phase. The equilibrium pressure in a CO₂ composition region larger than $y_{CO2} = 0.989$ is higher than that of the simple CO₂ hydrate systems. The inset in **Figure II-4-2** indicates the schematic phase diagram in the high CO₂ composition region, which is deduced from the results in the HFC-32+HFC-134a mixed hydrate (**Chapter II Section 3**). Although similar behavior was observed in the HFC-32+HFC-134a mixed-gas hydrate system (**Chapter II Section 3**), the isothermal phase equilibrium relations at T = 279.15 K in the CO₂+HFC-134a mixed-gas hydrate system do not exhibit a negative azeotropic-like behavior as observed in the HFC-32+HFC-134a mixed-gas hydrate system.



Figure II-4-2. Isothermal phase equilibria for the CO_2 +HFC-134a mixed-gas hydrate system at T = 279.15 K. The inset shows the schematic magnified-diagram for the right edge around the structural phase transition point.

p / MPa	X _{CO2}	УСО2	ZCO2
0.17	0.000	0.000	0.000
0.17	0.062	0.046	
0.18	0.127	0.106	
0.19	0.206	0.164	
0.20	0.273	0.222	
0.22	0.392	0.286	
0.23	0.540	0.386	
0.30			0.431
0.33	0.717	0.629	
0.49	0.827	0.790	
0.53			0.523
0.71	0.882	0.888	
0.95	0.940	0.932	
1.40			0.701
1.52	0.985	0.968	
1.94	0.990	0.973	
2.11	0.992	0.985	
2.47			0.713
2.49	0.995	0.985	
2.51	0.994	0.989	
2.53	0.995	0.989	
2.54	0.997	0.989	
2.56	0.998	0.991	
2.52	1.000	1.000	1.000

Table II-4-2. Isothermal phase equilibrium data in the CO₂+HFC-134a mixed-gas hydrate system at $T = 279.15 \text{ K.}^{a}$

^{*a*} u(p) = 0.01 MPa and $u(x_{CO2}) = u(y_{CO2}) = u(z_{CO2}) = 0.005$.

Figure II-4-3 shows typical Raman spectra of the CO₂+HFC-134a mixed-gas hydrate system at T = 279.15 K. The resonance peaks derived from the intramolecular vibration modes of the enclathrated CO₂ molecule were detected at $\Delta \nu = 1273$ cm⁻¹ and 1381 cm⁻¹ in the composition ranges of $z_{CO2} = 0.50-0.72$, while the peaks were detected at $\Delta v = 1275$ cm⁻¹ and 1381 cm⁻¹ in the composition region of $z_{CO2} \approx 0.99$. It has been reported that the Raman peak of CO₂ around $\Delta \nu =$ 1270 cm⁻¹ depends on the hydrate structures of s-I and s-II. The Raman shifts of $\Delta \nu = 1273$ cm⁻¹ and 1381 cm⁻¹ correspond to the CO₂ molecule encaged in the s-II hydrate [2]. The Raman shifts of $\Delta v = 1275 \text{ cm}^{-1}$ and 1381 cm⁻¹ agree well with the CO₂ molecule encaged in the s-I hydrates [1,2,4– 6]. The Raman peaks derived from the intramolecular C-H symmetric and C-H antisymmetric stretching vibration modes of the enclathrated HFC-134a molecule were detected at $\Delta v = 2977$ cm⁻¹ and 3010 cm⁻¹ in the composition ranges of $z_{CO2} = 0.50-0.72$. These shifts correspond to the HFC-134a molecule encaged in the L-cage of s-II hydrate (Chapter II Section 3). Moreover, the weak peak derived from the intramolecular C-H symmetric stretching vibration mode was detected at $\Delta v = 2985$ cm⁻¹, while the peak derived from the intramolecular C–H antisymmetric stretching vibration mode was too weak to be detected at the composition of $z_{CO2} \approx 0.99$. The existence of the peak at $\Delta v = 2985$ cm⁻¹ reveals that HFC-134a molecule is encaged in the M-cage of s-I hydrate under the coexistence of CO₂ like the HFC-32+HFC-134a mixed-gas hydrate system (Chapter II Section 3).



Figure II-4-3. Raman spectra of the intramolecular vibration modes of the enclathrated CO₂ (open inverse triangles) and HFC-134a (closed inverse triangles) molecules in the CO₂+HFC-134a mixed-gas hydrate system at T = 279.15 K.
These results of isothermal phase equilibrium relations and Raman spectra in the CO₂+HFC-134a mixed-gas hydrate system indicate that the structural phase transition occurs and a quadruple point (H_IH_{II}L₁G) exists at $p \approx 2.56$ MPa and $y_{CO2} \approx 0.991$; that is, this system exhibits the heterogeneous azeotropic-like behavior like the HFC-32+HFC-134a mixed-gas hydrate system.

CO₂+HFC-152a Mixed-Gas Hydrate System

Before measuring the phase equilibria in the CO₂+HFC-152a mixed-gas hydrate system, the cage occupancy of HFC-152a in the simple HFC-152a hydrate was investigated. The structure of HFC-152a hydrate remains s-I at pressures up to p = 370 MPa. The HFC-152a molecules occupy only M-cages of s-I simple HFC-152a hydrate in the presence of completely vacant small cages at pressures up to p = 370 MPa. Pressure dependence of the Raman shifts reveals that the HFC-152a molecule would be the largest s-I hydrate former.

Isothermal phase equilibrium relations for the CO₂+HFC-152a mixed-gas hydrate system at T = 279.15 K are shown in **Figure II-4-4** and summarized in **Table II-4-3**. The equilibrium CO₂ compositions in hydrate, aqueous, and gas phases increase in that order at a given pressure. The equilibrium pressure increases monotonically with an increasing of equilibrium CO₂ composition in all phases. The continuity in each equilibrium curve reveals that, unlike the CO₂+*c*-C₃H₆ mixed-gas hydrate system, no structural phase transition occurs in the CO₂+HFC-152a mixed-gas hydrate system at T = 279.15 K.



Figure II-4-4. Isothermal phase equilibria for the CO_2 +HFC-152a mixed-gas hydrate system at T = 279.15 K.

<i>p</i> / MPa	XCO2	УСО2	ZCO2
0.13	0.000	0.000	0.000
0.16	0.066	0.148	
0.20	0.233	0.327	
0.25	0.338	0.475	
0.30	0.435	0.562	
0.34			0.202
0.35	0.475	0.643	
0.44	0.563	0.711	
0.60	0.699	0.807	
0.76			0.360
0.79	0.781	0.864	
0.98	0.837	0.898	
1.14			0.525
1.19	0.906	0.924	
1.42	0.935	0.945	
1.66			0.763
1.60	0.956	0.958	
1.83	0.968	0.968	
2.09	0.987	0.981	
2.15			0.883
2.26	0.995	0.990	
2.33	0.998	0.992	
2.52	1.000	1.000	1.000

Table II-4-3. Isothermal phase equilibrium data in the CO₂+HFC-152a mixed-gas hydrate system at $T = 279.15 \text{ K.}^{a}$

^{*a*} u(p) = 0.01 MPa and $u(x_{CO2}) = u(y_{CO2}) = u(z_{CO2}) = 0.005$.

Figure II-4-5 shows typical Raman spectra of the CO₂+HFC-152a mixed-gas hydrate system at T = 279.15 K. The double Raman peaks derived from the intramolecular vibration modes of the enclathrated CO₂ molecule were detected at $\Delta \nu = 1275$ cm⁻¹ and 1381 cm⁻¹ in the whole composition range. The Raman shifts agree well with the CO₂ molecule encaged in the s-I hydrates [1,2,4–6]. The Raman peaks derived from the intramolecular CH₂ bending, C–C stretching, and CH₃ symmetric deformation vibrations of the enclathrated HFC-152a molecule were detected at $\Delta \nu$ = 1358 cm⁻¹, 1409 cm⁻¹, and 1446 cm⁻¹, respectively. The Raman shifts are in good agreement with that of the M-cage of the s-I simple HFC-152a hydrate. It is revealed that the crystal structure of the CO₂+HFC-152a mixed-gas hydrate remains s-I in the whole composition range at T = 279.15 K.



Figure II-4-5. Raman spectra of the intramolecular vibration modes of the enclathrated CO₂ (open inverse triangles) and HFC-152a (closed inverse triangles) molecules in the CO₂+HFC-152a mixed-gas hydrate system at T = 279.15 K.

II.4.5. Conclusions

The isothermal pressure-composition relations in the CO₂+HFC-134a and CO₂+HFC-152a mixed-gas hydrate systems were investigated under the three-phase equilibrium conditions at T = 279.15 K. The crystal structures of hydrates were analyzed based on the *in situ* Raman spectra. The CO₂+HFC-134a mixed-gas hydrate system exhibits the heterogeneous azeotropic-like behavior at $p \approx 2.56$ MPa and $y_{CO2} \approx 0.991$, while the CO₂+HFC-152a mixed-gas hydrate system shows straightforward equilibrium curves in the whole composition range.

II.4.6. Nomenclatures

ν	Wavenumber [m ⁻¹]
р	Pressure [Pa]
Т	Temperature [K]
и	Standard uncertainties of pressure [Pa], temperature [K], and mole fraction [-]
XCO2	Mole fraction of CO ₂ in aqueous phase (water free) [-]
YCO2	Mole fraction of CO ₂ in gas phase (water free) [-]
ZCO2	Mole fraction of CO ₂ in hydrate phase (water free) [-]
G	Gas phase
Н	Hydrate phase
HI	s-I hydrate phase
HII	s-II hydrate phase
L_1	Aqueous phase

II.4.7. References

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Section 5

Isothermal Phase Equilibria for the Xenon+Cyclopropane Mixed-Gas Hydrate System

II.5.1. Abstract

Isothermal three-phase equilibria of gas, aqueous, and hydrate phases for the Xe+c-C₃H₆ mixed-gas hydrate system were measured at T = 279.15 K and 289.15 K. The structural phase transitions from s-I to s-II and back to s-I, depending on the mole fraction of guest mixtures, occur in the Xe+c-C₃H₆ mixed-gas hydrate system. The isothermal pressure-composition relations have two local pressure minima. The most important characteristic in the Xe+c-C₃H₆ mixed-gas hydrate system relations exhibit the complex phase behavior involving two structural phase transitions and two homogeneous negative azeotropes. One of two structural phase transitions exhibits the heterogeneous azeotropic-like behavior.

II.5.2. Introduction

The structure of mixed-gas hydrates also depends on the composition of guest mixtures in addition to the system pressure and temperature. One of the most characteristic phenomena in the mixed-gas hydrate system is the structural phase transition from s-I to s-II and back to s-I, dependent on the composition of guest mixtures, while each guest species generates s-I hydrates. This phenomenon was first reported by Cailletet and Bordet [1] approximately 100 years ago. In 2000, Subramanian *et al.* [2] reported that the structural phase transition occurs in the CH₄+C₂H₆ mixed-gas hydrate system. The mechanism of structural phase transition is not yet clear. Therefore, it is very important to accumulate the thermodynamic properties of several mixed-gas hydrate systems in which the structural phase transition exists. The structural phase transition occurs in the CH₄+C₂H₆, CH₄+c-C₃H₆, CH₄+CF₄, and CO₂+c-C₃H₆ mixed-gas hydrate systems [2–

5], while it does not occur in the CH₄+C₂H₄, CH₄+CHF₃, CO₂+C₂H₆, and CO₂+CF₄ mixed-gas hydrate systems [6–8].

The Xe-containing mixed-gas hydrate systems have a possibility of the structural phase transition because van der Waals diameter of Xe (0.43 nm) is between those of CH₄ (0.41 nm) and CO₂ (0.50 nm). However, the investigation on the structural phase transition in Xe-containing mixed-gas hydrate systems is so rare. On the other hand, there are some reports on the phase equilibria of c-C₃H₆-containing mixed gas hydrates. The structural phase transition occurs in both CH₄+C₂H₆ and CH₄+c-C₃H₆ mixed-gas hydrate systems. The CH₄+c-C₃H₆ mixed-gas hydrate system [3] has a wider composition range of s-II hydrate than the CH₄+C₂H₆ and CH₄+CF₄ mixed-gas hydrate systems under isothermal conditions [2,4]. A structural phase transition occurs in the CO₂+c-C₃H₆ mixed-gas hydrate system [5], while it does not occur in the CO₂+C₂H₆ mixed-gas hydrate system [8]. These results imply that the Xe+c-C₃H₆ mixed-gas hydrate has a significant possibility of a structural phase transition, which is the reason I have focused on the Xe+c-C₃H₆ mixed-gas hydrate system.

In this section, isothermal phase equilibria for the Xe+c-C₃H₆ mixed-gas hydrate system were investigated under the three-phase (HL₁G) equilibrium conditions at T = 279.15 K and 289.15 K, where Xe and c-C₃H₆ generate the s-I hydrate under the present experimental conditions [9]. To clarify the structure of mixed-gas hydrates, single crystals of hydrates with various compositions were analyzed by means of *in situ* Raman spectroscopy.

II.5.3. Experimental

Materials

The materials used in the present study are summarized in **Table II-5-1**. All of them were used without further purification.

Material name	Source	Mole fraction purity
Xe	Air Water Inc.	> 0.99995
<i>c</i> -C ₃ H ₆	Takachiho Trading Co., Ltd.	> 0.995
Distilled water	Wako Pure Chemicals Ind., Ltd.	> 0.9999

 Table II-5-1. Information on the materials used in the present study.

Apparatus

The experimental apparatus for isothermal phase equilibrium measurement and Raman spectroscopy used in the present study are the same as described in **Chapter II Section 3**.

Procedure

The experimental procedures for isothermal phase equilibrium measurement and Raman spectroscopy performed in the present study are the same as described in **Chapter II Section 3**.

II.5.4. Results and Discussion

Three-phase equilibrium relations for the $Xe+c-C_3H_6$ mixed-gas hydrate system at T = 279.15K and 289.15 K are shown in Figures II-5-1 and 2, respectively. Three-phase equilibrium data for the Xe+c-C₃H₆ mixed-gas hydrate system are summarized in Tables II-5-2 and 3. Table II-5-2 summarizes the pressure (p)-composition (x_{Xe} , y_{Xe}) relations for each isotherm obtained without the dissociation of hydrate samples. Table II-5-3 summarizes the pressure (p)-composition (z_{Xe}) relation for each isotherm. The pressure-composition diagrams exhibit intensely curious behavior compared with those of various mixed-gas hydrate systems. At both temperatures, two discontinuous changes and two local pressure minima appear on each isotherm. The former indicate the existence of the structural phase transitions, which is supported by the Raman spectra mentioned below. One of two structural phase transitions, the structural phase transition between two local pressure minima, exhibits the heterogeneous azeotropic-like behavior. The other at the Xe-rich side exhibits the ordinary structural phase transition as reported in the CH₄+C₂H₆ mixed-gas hydrate system [2]. The local pressure minima and structural phase transition points in the $Xe+c-C_3H_6$ mixed-gas hydrate system are summarized in Table II-5-4. Considering the isothermal phase equilibria with the Raman spectra (mentioned below), the thermodynamically stable region of the s-II Xe+c-C₃H₆ mixed-gas hydrate is $y_{Xe} = 0.34-0.95$, $z_{Xe} = 0.40-0.74$ at T = 279.15 K and $y_{Xe} = 0.40-0.74$ 0.50–0.92, $z_{Xe} = 0.55$ –0.71 at T = 289.15 K. The s-II hydrate region becomes narrower with increase in the temperature.



Figure II-5-1. Isothermal phase equilibria for the $Xe+c-C_3H_6$ mixed-gas hydrate system at T = 279.15 K.



Figure II-5-2. Isothermal phase equilibria for the Xe+c-C₃H₆ mixed-gas hydrate system at T = 289.15 K.

<i>p</i> / MPa ^a	<i>y</i> Xe ^{<i>a</i>}	<i>X</i> Xe ^{<i>a</i>}	<i>p</i> / MPa ^a	<i>y</i> Xe ^{<i>a</i>}	<i>X</i> Xe ^{<i>a</i>}
	$T = 279.15 \text{ K}^{a}$			$T = 289.15 \text{ K}^{a}$	
0.153	0.000	0.000	0.550	0.000	0.000
0.148	0.038	0.007	0.530	0.015	0.001
0.144	0.089	0.021	0.472	0.090	0.039
0.143	0.159	_b	0.460	0.164	0.061
0.144	0.236	0.072	0.456	0.204	0.075
0.147	0.293	0.101	0.455	0.236	0.097
0.151	0.349	_b	0.457	0.277	0.124
0.150	0.373	_b	0.462	0.332	0.149
0.148	0.402	0.181	0.467	0.380	0.172
0.147	0.414	0.203	0.479	0.424	0.198
0.146	0.494	0.225	0.490	0.462	0.239
0.145	0.550	0.307	0.500	0.513	0.262
0.147	0.645	0.357	0.490	0.566	0.316
0.153	0.735	0.443	0.488	0.613	0.340
0.160	0.746	0.541	0.489	0.656	0.382
0.164	0.765	0.587	0.495	0.693	0.413
0.176	0.827	0.617	0.506	0.754	0.476
0.180	0.841	0.694	0.520	0.784	0.538
0.187	0.864	0.703	0.560	0.839	0.703
0.201	0.886	0.851	0.632	0.901	0.802
0.233	0.930	0.920	0.755	0.972	_b
0.262	0.954	0.948	0.774	1.000	1.000
0.273	0.973	0.969			
0.284	1.000	1.000			

Table II-5-2. Isothermal phase equilibrium data of gas and aqueous phases coexisting with $Xe+c-C_3H_6$ mixed-gas hydrates.

^{*a*} u(p) = 0.004 MPa, u(T) = 0.02 K, and $u(y_{Xe}) = u(x_{Xe}) = 0.005$.

^{*b*} The only p- y_{Xe} relation is listed in spite of the lack of x_{Xe} because the datum set is still meaningful for understanding the phase behavior.

<i>p</i> / MPa ^a	ZXe ^a	<i>p</i> / MPa ^a	Z.Xe ^a
T = 279	9.15 K ^a	T = 289	.15 K ^a
0.15	0.000	0.55	0.000
0.15	0.044	0.47	0.195
0.14	0.123	0.48	0.286
0.14	0.179	0.49	0.585
0.15	0.409	0.50	0.620
0.15	0.265	0.54	0.681
0.15	0.497	0.68	0.693
0.18	0.642	0.71	0.771
0.20	0.654	0.77	1.000
0.23	0.688		
0.26	0.868		
0.27	0.950		
0.28	1.000		

Table II-5-3. Isothermal pressure-composition relations in hydrate phase for the Xe+c-C₃H₆ mixed-gas hydrate system at T = 279.15 K and 289.15 K.

^{*a*} u(p) = 0.02 MPa, u(T) = 0.02 K, and $u(z_{Xe}) = 0.01$.

Table II-5-4. Summary of the local pressure minima and the structural phase transition points in the $Xe+c-C_3H_6$ mixed-gas hydrate system at T = 279.15 K and 289.15 K.

T / K^a		Local pressure minima		Structural phase	transition points
		s-I	s-II		
279.15	<i>X</i> Xe ^{<i>b</i>}	0.05 ± 0.02	0.30 ± 0.02	0.13 ± 0.02	0.93 ± 0.02
	yxe ^b	0.16 ± 0.01	0.55 ± 0.01	0.34 ± 0.01	0.95 ± 0.01
	ZXe ^b	0.16 ± 0.01	0.55 ± 0.01	0.27 ± 0.02 (s-I)	0.74 ± 0.02 (s-II)
				0.40 ± 0.02 (s-II)	0.86 ± 0.03 (s-I)
	<i>p</i> / MPa	0.143 ± 0.004	0.145 ± 0.004	0.152 ± 0.004	0.25 ± 0.01
289.15	x_{Xe}^{b}	0.10 ± 0.02	0.34 ± 0.02	0.26 ± 0.02	0.85 ± 0.02
	yxe ^b	0.23 ± 0.01	0.61 ± 0.01	0.50 ± 0.01	0.92 ± 0.01
	ZXe ^b	0.23 ± 0.01	0.61 ± 0.01	0.35 ± 0.05 (s-I)	0.71 ± 0.02 (s-II)
				0.55 ± 0.03 (s-II)	0.77 ± 0.02 (s-I)
	p / MPa	0.455 ± 0.004	0.488 ± 0.004	0.505 ± 0.004	0.70 ± 0.01

 $^{a}u(T) = 0.02$ K.

^b Due to the determinations by the extrapolation of isotherms.

The single local pressure minimum derived from the homogeneous negative azeotropic-like behavior has been reported in some mixed-gas hydrate systems [6,10,11]. The same behavior was observed in the HFC-32+HFC-134a mixed-gas hydrate system as mentioned in Chapter II Section 3. The existence of two local pressure minima on an isotherm is very rare. Holder and Manganiello [10] and Thakore and Holder [11] have stated that the negative azeotropic-like behavior is due to the competing effect of small and large guest molecules. The compositions at the local pressure minima of the s-I Xe+c-C₃H₆ mixed-gas hydrate are $y_{Xe} = z_{Xe} = 0.16 \pm 0.01$ at T = 279.15 K and y_{Xe} $z_{Xe} = 0.23 \pm 0.01$ at T = 289.15 K. In the case of the s-II Xe+c-C₃H₆ mixed-gas hydrate, the compositions are $y_{Xe} = z_{Xe} = 0.55 \pm 0.01$ at T = 279.15 K and $y_{Xe} = z_{Xe} = 0.61 \pm 0.01$ at T = 289.15K. If the Xe and c-C₃H₆ molecules occupy the small and large cages of both hydrate structures respectively, the ideal compositions, which are equal to the fractions of S-cage in the unit lattices, are $z_{Xe} = 0.25$ (s-I) and $z_{Xe} = 0.67$ (s-II). With increase in the temperature, the compositions at the local pressure minima are getting close to the fraction of S-cage in each hydrate structure and the negative azeotropic-like behavior becomes more remarkable. These results imply that the negative azeotropic-like behavior would result from both the compartmental occupancy of a suitable size of small and large guest species and the S-cage occupancy ratio of small guest species. Therefore, this discussion holds also in the HFC-32+HFC-134a mixed hydrate system mentioned in Chapter II Section 3, HFC-32 and HFC-134a could compartmentally occupy S-cages and L-cages of s-II hydrate at a pressure minimum, respectively.

As mentioned above, two discontinuous changes indicate the existence of the structural phase transitions. I confirmed this based on the Raman spectra corresponding to the intramolecular vibration modes of c-C₃H₆ in Xe+c-C₃H₆ mixed-gas hydrates with various compositions. Raman spectra corresponding to the ring-breathing and C–H stretching vibration modes of c-C₃H₆ in the Xe +c-C₃H₆ mixed-gas hydrates at T = 279.15 K and 289.15 K were measured. The results at T = 289.15 K have the same trend as those at T = 279.15 K. As typical spectra, the Raman spectra at T = 279.15 K are shown in **Figure II-5-3**. At $y_{Xe} = 0.34$ and 0.97, the Raman peak corresponding to the ring-breathing vibration mode of the c-C₃H₆ molecule is detected at $\Delta \nu = 1191$ cm⁻¹. On the other hand, at $y_{Xe} = 0.75$, 0.84, and 0.89, the Raman shift is $\Delta \nu = 1184$ cm⁻¹. The Raman shifts of $\Delta \nu = 1191$ cm⁻¹ and 1184 cm⁻¹ are in good agreement with those of c-C₃H₆ in the both s-I and s-II CH₄+c-C₃H₆ (and also CO₂+c-C₃H₆) mixed-gas hydrate systems, respectively [3,5]. Also, the peak shape and shift of the Raman spectra corresponding to the C–H stretching vibration mode are in good agreement with those of c-C₃H₆ in the both s-I and s-II cm⁻¹ (shown by the dotted lines) is derived from the c-C₃H₆ molecule in M-cages of s-I hydrate. Those at $\Delta \nu = 3009$ cm⁻¹ and 3024 cm⁻¹ (shown by the broken lines) are

derived from the c-C₃H₆ molecule in L-cages of s-II hydrate. These results prove the existence of structural phase transitions from s-I to s-II and back to s-I.



Figure II-5-3. Raman spectra corresponding to the ring-breathing ($\Delta \nu = 1190 \text{ cm}^{-1}$) and C–H stretching vibration ($\Delta \nu = 3000-3040 \text{ cm}^{-1}$) modes of *c*-C₃H₆ in the Xe+*c*-C₃H₆ mixed-gas hydrate at *T* = 279.15 K. The dotted and broken lines correspond to the Raman shifts in the s-I and s-II Xe +*c*-C₃H₆ mixed-gas hydrates, respectively.

II.5.5. Conclusions

Isothermal phase equilibria and Raman spectra in the $Xe+c-C_3H_6$ mixed-gas hydrate system were measured at T = 279.15 K and 289.15 K. The following important findings are obtained in the present study.

(1) Two kinds of homogeneous negative azeotropic-like behavior appear on an isotherm at the temperatures. The compositions of the local pressure minima are located at the vicinity of the cage fraction of the s-I and s-II hydrates.

(2) The phase behavior and the Raman spectra reveal the existence of two structural phase transitions from s-I to s-II and back to s-I. One of them exhibits the heterogeneous azeotropic-like behavior.

II.5.6. Nomenclatures

ν	Wavenumber [m ⁻¹]
р	Pressure [Pa]
Т	Temperature [K]
и	Standard uncertainties of pressure [Pa], temperature [K], and mole fraction [-]
XXe	Mole fraction of Xe in aqueous phase (water free) [-]
УХе	Mole fraction of Xe in gas phase (water free) [-]
ZXe	Mole fraction of Xe in hydrate phase (water free) [-]
G	Gas phase
Н	Hydrate phase
L_1	Aqueous phase

II.5.7. References

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Chapter III

Compartmental-Type Mixed Hydrates

Section 1

Phase Equilibrium Relations for Binary Mixed Hydrate Systems Composed of Carbon Dioxide and Cyclopentane Derivatives

III.1.1. Abstract

Thermodynamic stability boundaries of $CO_2+c-C_5H_{10}$, $CO_2+cyclopentanone$ (*c*-C₅H₈O), and $CO_2+fluorocyclopentane$ (*c*-C₅H₉F) mixed hydrates have been investigated at *T* = 274.19–293.72 K and up to *p* = 4.88 MPa. The phase equilibrium curves of the s-II mixed hydrates are located at temperatures remarkably higher than that of the s-I simple CO_2 hydrate. The highest temperature point of each four-phase (HL₁L₃G) equilibrium curve is *T* = 292.60 K (CO₂+*c*-C₅H₁₀ mixed hydrate system), 284.78 K (CO₂+*c*-C₅H₈O mixed hydrate system), and 293.72 K (CO₂+*c*-C₅H₉F mixed hydrate system). CO₂+bromocyclopentane (*c*-C₅H₉Br)+water system has also been investigated in the similar pressure and temperature range. However, the *c*-C₅H₉Br molecule is not enclathrated into any cage in the presence of CO₂.

III.1.2. Introduction

Hydrate is suggested to be utilized for some applications such as gas storage and transportation, gas isolation, gas separation, refrigeration, and so on. However, high pressures and low temperatures are naturally required for its formation and stabilization. Wherever hydrate is operated practically, therefore, it is necessary to moderate the working conditions. A reasonable way to moderate these conditions is the addition of the second guest species which is called a thermodynamic promoter [1–6].

In this section, four five-membered cyclic hydrocarbon molecules (c-C₅H₁₀, c-C₅H₈O, c-C₅H₉F, and c-C₅H₉Br) were selected as candidates for thermodynamic promoters. Five-membered cyclic hydrocarbons have a suitable molecular size for the L-cage occupancy of s-II hydrates, and

they may work as thermodynamic promoters [2–6]. The pressure-temperature relations for $CO_2+c-C_5H_{10}$ derivative+water systems were measured under four-phase equilibrium conditions (HL₁L₃G). The crystal structure of hydrate was analyzed by means of PXRD. Furthermore, the dissociation enthalpies of the $CO_2+c-C_5H_{10}$ derivative mixed hydrates were roughly estimated with Clapeyron equation under the assumption of full compartmental occupancy in S- and L-cages.

III.1.3. Experimental

Materials

The materials used in the present study are summarized in **Table III-1-1**. All of them were used without further purification.

Chemical name	Chemical name Source	
CO ₂	Neriki Gas Co., Ltd.	> 0.9999
<i>c</i> -C ₅ H ₁₀	Tokyo Chemical Ind. Co., Ltd.	> 0.98
$c-C_5H_8O$	Merck, Ltd.	> 0.99
c-C5H9F	SynQuest Laboratories, Inc.	> 0.9869
c-C5H9Br	Merck, Ltd.	> 0.99
Distilled water	Wako Pure Chemical Ind., Ltd.	> 0.9999

Table III-1-1. Information on the chemicals used in the present study.

<u>Apparatus</u>

The experimental apparatus for phase equilibrium measurement used in the present study is the same as described in **Chapter II Section 3** and shown in **Figure II-3-1**. The PXRD pattern was measured using a diffractometer (Rigaku, Ultima IV) with a Rigaku D/teX ultra high-speed position sensitive detector and CuK α X-ray (generation power: 40 kV, 50 mA).

Procedure

For phase equilibrium measurement, a desired volume of distilled water and c-C₅H₁₀ derivative was introduced into the evacuated high-pressure cell. The contents were pressurized with CO₂ up to a desired pressure. The cell was immersed into a thermostatic cooling bath. The contents were cooled and agitated to generate mixed hydrates. A magnetic stirrer was moved up and down

for agitation. After that, the temperature was gradually increased and decreased respectively to establish four-phase coexistence condition. When the pressure change was within 0.01 MPa and four-phase coexistence was confirmed visually, I determined that the system reached a four-phase equilibrium. Phase equilibrium curves stand vertically or rather retrograde at a pressure above the maximum temperature point. Temperature was increased very gradually (0.1 K every 1 h) to allow the amount of hydrate to decrease and the system to reach equilibrium at each step. The temperature and pressure where negligible amount of gas hydrates exists were determined as a four-phase equilibrium condition.

For PXRD, the structure of the CO₂+*c*-C₅H₈O mixed hydrate has not been reported. After the CO₂+*c*-C₅H₈O mixed hydrates were prepared in the high-pressure cell, the contents were pressurized with additional CO₂ so that almost all of free water was converted to mixed hydrate. The hydrate samples were taken from the cell at T = 263 K and kept at T = 77 K. Samples were ground and put flatly on the sample stage of the diffractometer. PXRD was performed at T = 143 K. The measurements were performed in the step scan mode with scan rate of 10° min⁻¹ and step size of 0.02°. The PXRD pattern indexing and cell refinement were performed with PowderX [7] and Chekcell [8] programs.

III.1.4. Results and Discussion

The four-phase (HL₁L₃G) equilibrium curves of the CO₂+*c*-C₅H₁₀ derivative mixed hydrate systems are shown in **Figure III-1-1** accompanied with the three kinds of three-phase (HL₁G, HL₂G, and L₁L₂G) equilibrium curves of the simple CO₂ hydrate system [9]. **Table III-1-2** summarizes the four-phase equilibrium relations of the CO₂+*c*-C₅H₁₀ derivative+water systems containing hydrates. The present four-phase (HL₁L₃G) equilibrium curve of the CO₂+*c*-C₅H₁₀ mixed hydrate system below p = 3 MPa agrees well with the reported data [10,11]. The dissociation temperatures of the present mixed hydrates except for CO₂+*c*-C₅H₉Br+water system are higher than that of the simple CO₂ hydrate at a certain pressure. On the other hand, the phase equilibrium curve (HL₁L₃G) of CO₂+*c*-C₅H₉Br+water system coincides with that of simple CO₂ hydrate system (HL₁G) [9]. It means that the *c*-C₅H₁₀, *c*-C₅H₈O, and *c*-C₅H₉F are enclathrated in hydrate cages. On the other hand, *c*-C₅H₉Br is not enclathrated in the presence of CO₂. Hereafter, CO₂+*c*-C₅H₉Br +water system is not mentioned. The enclathration of the *c*-C₅H₈O molecule was revealed in the present study. The largest pressure reduction from the equilibrium pressure of the simple CO₂ hydrate is observed in the CO₂+*c*-C₅H₉F hydrate system, followed by the CO₂+*c*-C₅H₁₀ and CO₂+*c*-C₅H₈O hydrate systems. Similar tendencies have been reported in the HFC-32+*c*-C₅H₁₀ and HFC-32+*c*-C₅H₁₀

C₅H₉F systems [5], the Kr+*c*-C₅H₁₀ and Kr+*c*-C₅H₉F systems [5], and fluoromethane (CH₃F)+*c*-C₅H₁₀ and CH₃F+*c*-C₅H₉F systems [6]. It is described in the literatures that *c*-C₅H₁₀ [12] and *c*-C₅H₉F [5,6] molecules occupy L-cages of s-II hydrates. PXRD pattern revealed that the crystal structure of the CO₂+*c*-C₅H₈O mixed hydrate is also s-II (*Fd3m*, $a = 1.7291 \pm 0.0010$ nm).



Figure III-1-1. Phase equilibrium relations of the CO₂+*c*-C₅H₁₀ derivative mixed hydrate systems.

T/\mathbf{K}^{a}	<i>p</i> / MPa ^{<i>a</i>}	T / K^a	<i>p</i> / MPa ^{<i>a</i>}	T / K^a	<i>p</i> / MPa ^{<i>a</i>}	T/K^{a}	<i>p</i> / MPa ^{<i>a</i>}
CO ₂ + <i>c</i> -C ₅ sys	H ₁₀ +water tem	CO ₂ + <i>c</i> -C ₅ sys	H ₈ O+water tem	CO ₂ + <i>c</i> -C ₅ sys	H9F+water tem	CO ₂ + <i>c</i> -C ₅ I sys	H9Br+water tem
280.16	0.08	274.19	0.47	283.18	0.05	274.29	1.41
280.76	0.11	274.49	0.48	284.21	0.13	275.08	1.56
281.22	0.15	275.42	0.56	285.20	0.22	276.21	1.78
283.17	0.34	276.46	0.66	286.22	0.33	277.16	1.99
284.04	0.45	277.36	0.77	287.24	0.46	278.26	2.27
285.11	0.60	278.39	0.91	288.24	0.61	279.19	2.58
286.09	0.76	279.36	1.07	289.23	0.81	280.20	2.94
287.07	0.96	280.34	1.26	290.23	1.04	281.13	3.35
288.03	1.20	281.34	1.50	291.25	1.36		
289.03	1.49	282.31	1.79	292.15	1.76		
290.03	1.85	283.24	2.17	293.19	2.46		
291.01	2.36	284.30	2.64	293.56	2.84		
291.21	2.45	284.78^{b}	3.21	293.72 ^b	3.21		
291.95	2.82	284.60	3.68	293.66	3.60		
292.45	3.31	293.47	4.09				
292.57	3.71	292.62	4.49				
292.60 ^b	4.10	291.89	4.79				
292.55	4.36						
292.29	4.71						
291.57	4.88						

Table III-1-2. Four-phase equilibrium conditions for $CO_2+c-C_5H_{10}$ derivative+water systems.

^{*a*} u(p) = 0.02 MPa and u(T) = 0.02 K.

_

^b The maximum equilibrium temperatures.

As the equilibrium pressure increases, the dissociation temperature monotonically increases at pressures below about 3 MPa. The equilibrium temperature scarcely changes at $p \approx 3$ MPa and the highest dissociation temperature is observed in the present mixed hydrate systems. The equilibrium temperature gradually decreases with an increase of pressure at pressures above the highest temperature point. The highest equilibrium temperature points in the CO₂+*c*-C₅H₈O, CO₂+*c*-C₅H₁₀, and CO₂+*c*-C₅H₉F mixed hydrate systems are located at p = 3.21 MPa and T = 284.78 K, p = 4.10 MPa and T = 292.60 K, and p = 3.21 MPa and T = 293.72 K, respectively. Based on Clapeyron equation (**Equation III-1-1**), the retrograde behavior was mainly caused by negative change of the molar volume (maybe derived from the partial molar volume of *c*-C₅H₁₀ derivative in L₃ phase as well as the slight change of S-cage occupancy of CO₂) during hydrate dissociation.

$$\frac{\mathrm{d}p}{\mathrm{d}T} = \frac{\Delta_{\mathrm{hyd}}H}{T\,\Delta_{\mathrm{hyd}}\nu}$$
(Equation III-1-1)

where $\Delta_{hyd}H$ and $\Delta_{hyd}v$ mean the enthalpy and molar volume changes through the hydrate dissociation, respectively.

The existence of the quintuple point (HL₁L₂L₃G) was reported in CO₂+tetrahydropyran, CO₂+cyclobutanone, CO₂+cyclohexane, and CO₂+methylcyclohexane mixed hydrate systems [14]. In these cases, five four-phase equilibrium curves (HL₁L₂L₃, HL₁L₂G, HL₁L₃G, HL₂L₃G, and L₁L₂L₃G) converge at the quintuple point (HL₁L₂L₃G) as reported elsewhere [14]. I have explored four-phase equilibrium curves other than HL₁L₃G by changing the ratio of materials. For the exploration of the HL₂L₃G or HL₁L₂G curves, the phase L₁ or L₃ needs to disappear after hydrate formation. A small amount of either water or *c*-C₅H₁₀ derivative was added into the normal volume (2 cm³) of *c*-C₅H₁₀ derivative or water, respectively. However, hydrates were not formed and then the phase condition was the three-phase coexistence of L₁L₃G in both experiments. Theoretically, a quintuple point must have five four-phase equilibrium curves. However, no other curve than HL₁L₃G), the coexistence point of both s-II mixed hydrate and s-I simple CO₂ hydrate would be placed near the quadruple point in simple CO₂ hydrate system (HL₁L₂G). It is because the present CO₂+*c*-C₅H₁₀ derivative mixed hydrates are s-II and simple CO₂ hydrate is s-I, whereas none of the *c*-C₅H₁₀ derivatives is able to occupy the M-cage of s-I hydrate.

The overall dissociation enthalpies of $CO_2+c-C_5H_{10}$ derivative mixed hydrates were estimated with Clapeyron equation (**Equation III-1-1**). The partial differentials of pressure with respect to temperature were obtained from the correlated four-phase equilibrium curves obtained in the present study. IUPAC recommended equations of state [15] and the correlation of Wagner and Pruss [16] were used to calculate the molar volumes of CO_2 and H_2O , respectively. The molar volumes of $c-C_5H_{10}$ derivatives were calculated from densities measured in the present study. The molar

volumes of the s-I and s-II hydrates were calculated from each lattice constant (a = 1.20 nm (s-I), a= 1.73 nm (s-II) [1]). I made some assumptions in the calculation of the overall dissociation enthalpy of the $CO_2+c-C_5H_{10}$ derivative mixed hydrates. CO_2 and $c-C_5H_{10}$ derivative compartmentally occupy S- and L-cages of the s-II hydrate (2CO₂·c-C₅H₁₀ derivative·17H₂O), respectively. In the case of the simple CO₂ hydrate, it was assumed that CO₂ molecules completely occupy the S-cages and M-cages of s-I hydrate (CO₂·5.75H₂O). It was considered that the pressure dependence on the molar volume for the $c-C_5H_{10}$ derivative was negligible and the lattice parameter of the s-II hydrate was constant. To minimize the errors caused by the pressure effect, I have estimated the dissociation enthalpy only in a low pressure range. The dissociation enthalpies per mole of hydrates were 308 ± 29 kJ mol⁻¹ (p = 0.60-0.96 MPa and T = 285.11-287.07 K), 202 ± 5 kJ mol⁻¹ (p = 0.66-0.91 MPa and T = 276.46-278.39 K), and 379 ± 25 kJ mol⁻¹ (p = 0.46-0.81 MPa and T = 287.24 - 289.23 K) in the CO₂+*c*-C₅H₁₀, CO₂+*c*-C₅H₈O, and CO₂+*c*-C₅H₉F mixed hydrates, respectively. The dissociation enthalpy of the CO₂+*c*-C₅H₉F mixed hydrate is the highest, followed by the $CO_2+c-C_5H_{10}$ and $CO_2+c-C_5H_8O$ mixed hydrates. The dissociation enthalpy of the simple CO_2 hydrate per mole of CO_2 , which is estimated with the same procedure, is approximately 68 ± 6 kJ mol⁻¹ (T = 275.37 - 281.21 K, p = 1.54 - 3.28 MPa) on the three-phase equilibrium curve (HL₁G) [9]. For comparison, the estimated dissociation enthalpies per mole of CO₂ are 154 ± 15 kJ mol⁻¹ (p = 0.60–0.96 MPa and T = 285.11–287.07 K), 101 \pm 2 kJ mol⁻¹ (p = 0.66–0.91 MPa and T = 276.46– 278.39 K), and 190 \pm 13 kJ mol⁻¹ (p = 0.46-0.81 MPa and T = 287.24-289.23 K) in the CO₂+c- C_5H_{10} , $CO_2+c-C_5H_8O$, and $CO_2+c-C_5H_9F$ mixed hydrates, respectively.

III.1.5. Conclusions

Phase equilibrium curves of the $CO_2+c-C_5H_{10}$ derivative+water systems have been investigated. *c*-C₅H₉F, *c*-C₅H₁₀, and *c*-C₅H₈O form s-II mixed hydrates with CO₂ and the equilibrium pressures of the mixed hydrates are lower than those of the simple CO₂ hydrate, whereas *c*-C₅H₉Br does not form hydrate in the presence of CO₂. The CO₂+*c*-C₅H₉F hydrate has the highest dissociation temperature followed by the CO₂+*c*-C₅H₁₀ hydrate and CO₂+*c*-C₅H₈O hydrate at a certain pressure. Except for the CO₂+*c*-C₅H₉Br hydrate system, each four-phase equilibrium curve has the maximum dissociation temperature at *p* ≈ 3 MPa, and that of the CO₂+*c*-C₅H₉F mixed hydrate system is the highest in the present mixed hydrate systems. Except for the CO₂+*c*-C₅H₉Br hydrate system, four-phase equilibrium curves (HL₁L₃G) show retrograde behavior and approach to the structural transition point between s-I and s-II at pressures above the maximum dissociation point. The dissociation enthalpies of the $CO_2+c-C_5H_9F$, $CO_2+c-C_5H_{10}$, and $CO_2+c-C_5H_8O$ mixed hydrates decrease in this order.

III.1.6. Nomenclatures

a	Lattice constant [m]
Н	Enthalpy [J mol ⁻¹]
р	Pressure [Pa]
Т	Temperature [K]
и	Standard uncertainties of pressure [Pa] and temperature [K]
v	Molar volume [m ³ mol ⁻¹]
G	Gas phase
Н	Hydrate phase
L_1	Aqueous phase
L_2	CO ₂ -rich liquid phase
L ₃	<i>c</i> -C ₅ H ₁₀ derivative-rich liquid phase
Q2	Quadruple point of hydrate, aqueous, CO2-rich liquid, and gas phases

III.1.7. References

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Section 2

Thermodynamic Stability Boundaries and Structures of Methane+Monohalogenated Cyclopentane Mixed Hydrates

III.2.1. Abstract

Thermodynamic stability boundaries of mixed hydrates composed of CH₄ and c-C₅H₉F, chlorocyclopentane (c-C₅H₉Cl), or c-C₅H₉Br were investigated at T = 274.17-306.19 K and pressures up to p = 9.93 MPa under four-phase (HL₁L₃G) equilibrium conditions. The each boundary is laid at pressures lower than that of simple CH₄ hydrate. Especially, the CH₄+c-C₅H₉F mixed hydrate system has the lowest equilibrium pressure among three mixed hydrate systems investigated in the present study. Also, crystal structure of CH₄+c-C₅H₉Cl and CH₄+c-C₅H₉Br mixed hydrates was determined by means of PXRD. The PXRD patterns reveal that there is the limit of L-cage occupancy between c-C₅H₉Cl and c-C₅H₉Br.

III.2.2. Introduction

Hydrate enclathrates large amount of gas such as H_2 , CH_4 , and CO_2 . Therefore, hydrate is suggested to be utilized for gas storage, transportation, and separation. However, high pressures and/or low temperatures are required for the formation and stabilization of hydrates, which is one of the most serious problems to be solved. Adding large guest species as a thermodynamic promoter to a simple gas hydrate system, the gas and large guest species occupy small and large cages respectively, resulting that s-II or s-H mixed hydrate can form at thermodynamic conditions milder than simple gas hydrate (i.e. lower pressure). That is why s-II and s-H hydrates have drawn attention in the field of natural gas storage and transportation. So far, many large guest species forming s-II and s-H hydrates were reported [1]. In **Chapter III Section 1**, I stated that $c-C_5H_{10}$, $c-C_5H_8O$, and $c-C_5H_9F$ form s-II hydrates with CO_2 . However, the occupancy limit for the L-cage of s-II hydrate former has not been clarified yet. Which cage large guest species occupy gives fundamentally significant information about the occupancy limits of large guest species for the Lcage of s-II hydrate.

In this section, three monohalogenated cyclopentanes (*c*-C₅H₉X, X = F, Cl, and Br) are adopted as large guest species. c-C₅H₁₀ and c-C₅H₉F form s-II hydrates and they occupy only Lcages [2–4]. It has not been reported whether c-C₅H₉Cl forms hydrates or not. According to previous reports mentioned above and the results in **Chapter III Section 1**, c-C₅H₉Cl is supposed to be either s-II or s-H hydrate former because it is larger than c-C₅H₉F and smaller than c-C₅H₉Br. Additionally, taking into consideration the previous studies on Kr+c-C₅H₉F [3], HFC-32+c-C₅H₉F [3], CH₃F+c-C₅H₉F [4], and CO₂+c-C₅H₉F mixed hydrates (**Chapter III Section 1**), CH₄+c-C₅H₉F mixed hydrate should be formed under drastically mild conditions. The four-phase (HL₁L₃G) equilibrium curves of the CH₄+c-C₅H₉F, CH₄+c-C₅H₉Cl, and CH₄+c-C₅H₉Br mixed hydrate systems were measured. The structure of the CH₄+c-C₅H₉Cl mixed hydrate, in addition, was clarified based on PXRD pattern.

III.2.3. Experimental

Materials

The materials used in the present study are summarized in **Table III-2-1**. All of them were used without further purification.

Chemical name	Source	Mole fraction purity
CH ₄	Liquid Gas Co., Ltd.	> 0.9999
c-C5H9F	SynQuest Laboratories, Inc.	> 0.9869
c-C5H9Cl	Wako Pure Chemical Ind., Ltd.	> 0.98
c-C5H9Br	Merck Ltd.	> 0.99
Distilled water	Wako Pure Chemical Ind., Ltd.	> 0.9999

 Table III-2-1. Information on the chemicals used in the present study.

<u>Apparatus</u>

The experimental apparatus for phase equilibrium measurement and PXRD used in the present study are the same as described in Chapter II Section 3 and Chapter III Section 1, respectively.

Procedure

The experimental procedures for phase equilibrium measurement and PXRD performed in the present study are the same as described in **Chapter III Section 1**.

III.2.4. Results and Discussion

The PXRD patterns of the CH₄₊*c*-C₅H₉Cl and CH₄₊*c*-C₅H₉Br mixed hydrates obtained in the present study are shown in **Figure III-2-1**. The typical pattern of s-II hydrate was observed in the CH₄₊*c*-C₅H₉Cl mixed hydrate. The space group and the lattice constant of the CH₄₊*c*-C₅H₉Cl mixed hydrate are *Fd3m* and $a = 1.734 \pm 0.003$ nm, respectively. On the other hand, the typical pattern of s-H hydrate was detected in the CH₄₊*c*-C₅H₉Br mixed hydrate. The space group and the lattice constants of the CH₄₊*c*-C₅H₉Br mixed hydrate are *P6/mmm* and $a = 1.232 \pm 0.001$ nm, $c = 1.019 \pm 0.001$ nm, respectively. The present study firstly proved that *c*-C₅H₉Cl is enclathrated into the L-cages of s-II hydrate in the presence of CH₄. One of the most surprising findings is that *c*-C₅H₉Br is enclathrated into the U-cages of s-H hydrate in the presence of CO₂ (**Chapter III Section 1**). This result indicates that CH₄ is more helpful than CO₂ for the enclathration of large guest species. The PXRD patterns also indicate that there is the limit of L-cage occupancy between *c*-C₅H₉Cl and *c*-C₅H₉Br, the former is one of the largest guest species that occupy the L-cages of s-H hydrate.



Figure III-2-1. PXRD patterns obtained from $CH_4+c-C_5H_9Cl$ and $CH_4+c-C_5H_9Br$ mixed hydrate samples prepared in the present study (recorded at T = 173 K).

Thermodynamic stability boundaries of $CH_4+c-C_5H_9X$ (X = F, Cl, and Br) mixed hydrates measured in the present study under four-phase (HL₁L₃G) equilibrium conditions are shown in Figure III-2-2 accompanied with those of the $CH_4+c-C_5H_{10}$ mixed hydrate (s-II) [5,6] and the simple CH₄ hydrate (s-I) [7]. The obtained four-phase equilibrium data are also listed in Table **III-2-2**. The data for the $CH_4+c-C_5H_9Br$ mixed hydrate agree well with the data recently reported in [8]. The four-phase equilibrium curves are located at a pressure side lower than the three-phase (HL₁G) equilibrium curve of s-I simple CH₄ hydrate system. All of the *c*-C₅H₉X adopted in the present study work as a thermodynamic promoter. CH₄+c-C₅H₉F mixed hydrate is the most thermodynamically stable in the present study, followed by CH₄+*c*-C₅H₉Cl and CH₄+*c*-C₅H₉Br mixed hydrates. Thermodynamic stability region of the s-II CH₄+c-C₅H₉F mixed hydrate is quite different from that of the s-II CH₄+c-C₅H₉Cl mixed hydrate. It implies that c-C₅H₉F has significantly appropriate molecular size and shape to occupy L-cage, whereas c-C₅H₉Cl may be a little bit too large for the L-cage occupancy. Especially, the CH₄+c-C₅H₉F hydrate is formed at thermodynamic conditions milder than $CH_4+c-C_5H_{10}$ hydrate [5,6], where $c-C_5H_{10}$ is a remarkable thermodynamic promoter forming s-II hydrates. The same tendency is observed in the Kr+c-C₅H₁₀ and Kr+c-C₅H₉F mixed hydrates [3], HFC-32+c-C₅H₁₀ and HFC-32+c-C₅H₉F mixed hydrates [3], $CH_3F+c-C_5H_{10}$ and $CH_3F+c-C_5H_9F$ mixed hydrate [4], and $CO_2+c-C_5H_{10}$ and $CO_2+c-C_5H_9F$ mixed hydrates (Chapter III Section 1).



Figure III-2-2. Four-phase (HL₁L₃G) equilibrium relations in the CH₄+c-C₅H₉X (X = F, Cl, and Br)+water ternary systems.

T / K^a	p / MPa ^a	T / K^a	p / MPa ^a	T / K^a	<i>p</i> / MPa ^a
$CH_4+c-C_5H_9H$	F+water system	CH ₄ + <i>c</i> -C ₅ H ₉ Cl+water system		CH ₄ + <i>c</i> -C ₅ H ₉ B	r+water system
283.17	0.03	274.24	1.11	274.17	1.82
284.17	0.08	274.37	1.13	275.15	2.03
285.12	0.14	275.24	1.28	276.19	2.32
286.12	0.21	276.11	1.46	277.13	2.61
287.21	0.29	276.98	1.65	278.12	2.93
288.18	0.38	277.48	1.77	279.15	3.30
289.17	0.49	278.74	2.11	280.05	3.65
290.16	0.60	279.73	2.42	281.12	4.15
291.14	0.74	280.59	2.71	282.10	4.68
292.14	0.90	281.20	2.98	282.61	5.00
293.12	1.07	281.91	3.29	283.18	5.45
294.14	1.28	282.68	3.68	284.12	6.03
295.13	1.52	283.47	4.09	285.11	6.87
296.10	1.79	284.55	4.72	286.14	7.92
297.09	2.10	284.15	4.54	287.15	9.08
298.19	2.51	285.14	5.17	287.74	9.93
299.16	2.92	286.16	5.99		
300.14	3.41	287.10	6.85		
301.11	3.98	288.19	8.01		
302.12	4.65	289.19	9.33		
303.11	5.44				
304.12	6.37				
305.10	7.46				
306.19	8.93				

Table III-2-2. Four-phase equilibrium conditions for $CH_4+c-C_5H_9X$ (X = F, Cl, and Br)+water systems.

 $^{a} u(p) = 0.02$ MPa and u(T) = 0.02 K.

III.2.5. Conclusions

Thermodynamic stabilities of CH₄+c-C₅H₉F, CH₄+c-C₅H₉Cl, and CH₄+c-C₅H₉Br mixed hydrate systems were investigated. In addition, the structure of these mixed hydrates was clarified. Compared to the s-I simple CH₄ hydrate, the s-II CH₄+c-C₅H₉F mixed hydrate is much more thermodynamically stable, followed by s-II CH₄+c-C₅H₉Cl, and s-H CH₄+c-C₅H₉Br mixed hydrates. c-C₅H₉F works as a thermodynamic promoter much greater than c-C₅H₉Cl and c-C₅H₉Br. CH₄ works as a help gas for the enclathration of c-C₅H₉Br, whereas CO₂ does not. There is the limit of L-cage occupancy between c-C₅H₉Cl and c-C₅H₉Br. Comprehensively speaking, c-C₅H₉F is one of the best thermodynamic promoters, which is helpful for hydrate utilization.

III.2.6. Nomenclatures

a	Lattice constant [m]
с	Lattice constant [m]
р	Pressure [Pa]
Т	Temperature [K]
и	Standard uncertainties of pressure [Pa] and temperature [K]
G	Gas phase
Н	Hydrate phase
L_1	Aqueous phase
L ₃	<i>c</i> -C ₅ H ₉ X-rich liquid phase

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Section 3

Extraordinary Stability of Structure-H Methane+1,1,2,2,3,3,4-Heptafluorocyclopentane Mixed Hydrate at Pressures up to 373 MPa

III.3.1. Abstract

Thermodynamic stability boundary of s-H hydrates with large guest species (thermodynamic promoters) and CH₄ at extremely high pressures has been almost unclear. In the present study, the four-phase equilibrium relations in the s-H CH₄+1,1,2,2,3,3,4-heptafluorocyclopentane (1,1,2,2,3,3,4-HFCP) mixed hydrate system were investigated at T = 281.05-330.12 K and pressures up to p = 373 MPa. The difference between equilibrium pressures in the s-H CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate system and the s-I simple CH₄ hydrate system gets larger with increase in temperature. The s-H CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate survives even at p = 373 MPa and T = 330 K without any structural phase transition. The maximum temperature where the s-H CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate is thermodynamically stable is likely to be beyond that of the s-H simple CH₄ hydrate.

III.3.2. Introduction

In general, simple gas hydrate belongs to either s-I or s-II hydrate, except at extremely high pressures [1–3]. In a part of s-II and s-H hydrates, small and large guest species occupy small and large cages, compartmentally. If the large guest species does not form hydrates without small guest species, the small guest species is called a help gas.

Recently, s-H hydrates have drawn much attention as a new medium for the natural gas transportation system [4]. The equilibrium pressure of an s-H hydrate formed by the addition of a small amount of large guest species is smaller than that of a simple gas hydrate (s-I or s-II, dependent on help gases). Furthermore, a larger amount of help gas can be stored because s-H

hydrates have the largest storage potential in small cage(s) among s-I, s-II, and s-H hydrates by stabilizing large cage with large guest species [4]. The thermodynamic properties and storage abilities of s-H hydrates depend on the chemical and physical properties of large guest species [5–7], for example, the molecular volume, the molar mass, the shape of large guest species, and so on.

One of the characteristic phenomena in an s-H hydrate system is the existence of the structural phase transition point, where an s-H mixed hydrate is collapsed and instead a simple gas hydrate, which contains only a help gas without large guest species, is formed. For example, in the case of the utilization of CH₄ as help gas, phase equilibrium curve in the s-H mixed hydrate systems comes close to that of the simple CH₄ hydrate system as pressures and temperatures rise. Finally, the curve of the s-H mixed hydrate systems crosses that of the simple CH₄ hydrate system at the structural phase transition point. For example, CH₄+cis-4-methylcyclohexanol [6], Xe+1,1dimethylcyclohexane (1,1-DMCH) [7], CO₂+2,2-dimethylbutane [8], CH₃F and +methylcyclohexane [9] mixed hydrate systems exhibit such behavior. In other s-H mixed hydrate systems where such structural phase transition has not been reported, in addition, it has been considered that the s-H mixed hydrate is collapsed and changed to simple CH₄ hydrate at pressures higher than tens of MPa. It is because the slope (dp/dT) of phase equilibrium curve of the typical s-H mixed hydrate is larger than that of the simple gas hydrate containing only the once-help gas.

There are some important studies on the phase equilibria for the simple CH₄ hydrate system at extremely high pressures. Dyadin *et al.* [1] have reported that the simple CH₄ hydrate has the structural phase transition point at p = 620 MPa and T = 320 K on the three-phase (HL₁G) equilibrium curve. They have claimed that phase equilibrium curve after the phase transition intersects the melting curve of ice-VI at p = 1.62-1.65 GPa and T = 331 K, while the new crystal lattice was not identified definitely. Loveday *et al.* [2] have reported that the simple CH₄ hydrate above p = 620 MPa is very similar to s-H hydrate based on the X-ray and neutron diffraction analyses. Moreover, Kumazaki *et al.* [3] have reported the Raman spectra in a single crystal of the s-H CH₄ hydrate formed under three-phase equilibrium conditions (p = 1.0 GPa, T = 323 K).

In order to elucidate the characteristics of s-H hydrates, it is very important to investigate the phase equilibrium relations in s-H CH₄+large guest species mixed hydrate systems at higher pressures than ever, especially several hundred MPa. In this section, CH₄ and 1,1,2,2,3,3,4-HFCP are adopted as a help gas and a large guest species, respectively. First, I identified the structure of CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate because it has not been reported whether 1,1,2,2,3,3,4-HFCP forms hydrates. Thermodynamic stability of the CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate was investigated at pressures up to p = 373 MPa.
III.3.3. Experimental

Materials

The materials used in the present study are summarized in **Table III-3-1**. All of them were used without further purification.

Chemical name	Source	Mole fraction purity
CH ₄	Liquid Gas Co., Ltd.	> 0.9999
1,1,2,2,3,3,4-HFCP	Zeon Corp.	> 0.98
Distilled water	Wako Pure Chemical Ind., Ltd.	> 0.9999

 Table III-3-1. Information on the chemicals used in the present study.

Apparatus

The experimental apparatus for phase equilibrium measurement and PXRD used in the present study are the same as described in Chapter II Section 3 and Chapter III Section 1, respectively.

Procedure

The experimental procedures for phase equilibrium measurement and PXRD performed in the present study are the same as described in **Chapter III Section 1**. The experimental procedure for Raman spectroscopy done in the present study is the same as described in **Chapter II Section 3**.

III.3.4. Results and Discussion

The PXRD pattern of the CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate prepared at p = 3.31 MPa and T = 282.44 K is shown in **Figure III-3-1**. The typical pattern derived from s-H hydrate (hexagonal, *P6/mmm*) was detected in the CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate. The lattice constants are $a = 1.221 \pm 0.002$ nm and $c = 1.002 \pm 0.001$ nm at T = 173 K.



Figure III-3-1. PXRD pattern of CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate prepared at p = 3.31 MPa and T = 282.44 K (recorded at T = 173 K).

Four-phase equilibrium relations of HL₁L₃G and S₃L₁L₃G in the CH₄+1,1,2,2,3,3,4-HFCP +water ternary system are shown in Figure III-3-2 and summarized in Table III-3-2. In the whole pressure and temperature range of the present study, the four-phase equilibrium curve (HL_1L_3G) is laid at a lower pressure side than the three-phase equilibrium curve (HL₁G) in simple CH₄ hydrate system. Until now, it had been believed that the s-H mixed hydrate is collapsed at the intersection between the four-phase equilibrium curve of s-H hydrates and the three-phase equilibrium curve of simple gas hydrates containing only the once-help gas in s-H hydrate system. In fact, such phase behavior has been reported in some s-H hydrates helped by CH_4 [6] and Xe [7]. Surprisingly, the four-phase equilibrium curve (HL₁L₃G) in the CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate system does not intersect with the three-phase equilibrium curve (HL_1G) in the simple CH_4 hydrate system up to p = 373 MPa and T = 330.12 K. Rather, the difference between both equilibrium curves gets larger with increases in temperature and pressure. The four-phase equilibrium curve of HL₁L₃G exhibits no structural phase transition up to p = 373 MPa and T = 330.12 K. In addition, there is no significant difference in Raman shifts of the intramolecular vibration modes of the CH₄ and 1,1,2,2,3,3,4-HFCP molecules except for the weak pressure and temperature dependences, as shown in Figure III-3-3. On the other hand, the structure of simple CH₄ hydrate changes from s-I to s-H at p = 620 MPa and T = 320 K on the three-phase equilibrium curve (HL₁G) [1–3]. In the s-H simple CH₄ hydrate system, multiple CH₄ molecules surely occupy U-cage [2,3,10]. Directly comparing the stability boundary of the s-H CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate with that of the s-H simple CH₄ hydrate, it is suggested that the contribution of large guest species to the stability of s-H hydrates is crucially significant.



Figure III-3-2. Four-phase (HL₁L₃G and S₃L₁L₃G) equilibrium relations in the CH₄+1,1,2,2,3,3,4-HFCP+water ternary system.

T / K^a	p / MPa a	T / K a	p / MPa a
HI	$_1L_3G$	S_3L_1	L ₃ G
281.19	2.85	287.73	1.35
281.61	3.00	285.58	1.81
282.34	3.26	282.98	2.37
283.17	3.64	281.99	2.60
283.94	4.02	281.16	2.77
284.81	4.44		
285.12	4.63		
285.40	4.77		
285.78	5.02		
286.61	5.56		
287.30	6.06		
288.06	6.69		
289.12	7.68		
290.16	8.76		
290.69	9.41		
290.90	9.68		
293.35	13.3		
296.96	22.9		
298.67	29.9		
301.13	42.2		
303.69	57.6		
305.89	71.6		
308.53	92.0		
312.90	140		
315.59	168		
318.45	203		
323.83	270		
326.00	301		
328.56	344		
330.12	373		

Table III-3-2. Four-phase equilibrium data in the CH₄+1,1,2,2,3,3,4-HFCP+water ternary system.

 $^{a} u(p) = 0.02$ MPa at p < 10 MPa, u(p) = 0.2 MPa at p = 10-100 MPa, u(p) = 2 MPa at p > 100 MPa, and u(T) = 0.02 K.



Figure III-3-3. Raman spectra of the intramolecular vibrations of the enclathrated CH₄ (open inverse triangle) and 1,1,2,2,3,3,4-HFCP (closed inverse triangle) molecules in the CH₄+1,1,2,2,3,3,4-HFCP mixed hydrates.

Among the large guest species forming s-H hydrates, 1,1-DMCH has been considered to be the most suitable for the s-H hydrate generation (that is, as a thermodynamic promoter) from the viewpoint of the equilibrium pressure reduction [13,14]. The equilibrium pressure of the CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate is slightly higher than that of CH₄+1,1-DMCH mixed hydrate at pressures up to p = 10 MPa. Therefore, 1,1,2,2,3,3,4-HFCP would also be one of the suitable large guest species for the s-H hydrate generation.

III.3.5. Conclusions

1,1,2,2,3,3,4-HFCP generates an s-H hydrate in the presence of CH₄. Thermodynamic stability of the s-H CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate was investigated at pressures up to p = 373 MPa. The most important finding is that the s-H CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate remains thermodynamically stable even at p = 373 MPa and T = 330 K without any structural phase transition, although it had been believed that s-H hydrates with help gases would be collapsed and transformed to the simple hydrates enclathrating only help gases. Moreover, the stability boundary of the s-H CH₄+1,1,2,2,3,3,4-HFCP mixed hydrate is located beside that of the s-H CH₄+1,1-DMCH mixed hydrate. 1,1,2,2,3,3,4-HFCP would be one of the most suitable large guest species for the equilibrium pressure reduction derived from s-H hydrate formation.

III.3.6. Nomenclatures

θ	Diffraction angle [°]
ν	Wavenumber [m ⁻¹]
a	Lattice constant [m]
с	Lattice constant [m]
р	Pressure [Pa]
Т	Temperature [K]
и	Standard uncertainties of pressure [Pa] and temperature [K]
G	Gas phase
Н	Hydrate phase
L_1	Aqueous phase
L ₃	1,1,2,2,3,3,4-HFCP-rich liquid phase
S ₃	Solid 1,1,2,2,3,3,4-HFCP phase

III.3.7. References

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Section 4

Structural Phase Transitions of Methane+Ethane Mixed-Gas Hydrate Induced by 1,1-Dimethylcyclohexane

III.4.1. Abstract

With the combination of natural gases and thermodynamic promoters in mind, $CH_4+C_2H_6+1,1-DMCH+$ water system was investigated by means of Raman spectroscopy and isothermal phase equilibrium measurements under four-phase (HL₁L₃G) equilibrium conditions at T = 288.15 K. The results suggest that three kinds of hydrate structures emerge at T = 288.15 K in the $CH_4+C_2H_6+1,1-DMCH+$ water system. The hydrate structure for this system changed from s-H to s-I via s-II with increase in the mole ratio of C₂H₆ to CH₄.

III.4.2. Introduction

The s-H hydrate formation needs both small guest species in the S-, S'-cages and large guest species in the U-cage, respectively (except for a hexagonal structure of simple hydrates at extreme pressures [1,2]). The former is called as a help-gas.

It has been reported that many kinds of molecules are encaged in hydrates, from small one (e.g. H₂) to large one (e.g. methylcyclohexane [3–6]). Gas hydrates have a large amount of potential for many fields. For example, H₂, CO₂, and CH₄ hydrates have drawn much attention to be used as gas storage medium [7]. However, these hydrates are generated at relatively high pressure and low temperature conditions, so it is necessary to moderate the conditions of hydrate formation for practical applications of gas hydrates. Generally speaking, the equilibrium pressure of the s-H hydrate containing help-gas and large guest species is lower than that of the simple hydrate (s-I or s-II) containing only a help-gas. The degree of the pressure drop depends on large guest species [8]. The s-H CH₄+1,1-DMCH hydrate exhibits quite large depression of the equilibrium pressure in the s-H hydrates containing CH₄ as help-gas [9].

In addition, the crystal structure depends on the guest concentration as reported in the $CH_4+C_2H_6$ [10–13], $CH_4+c-C_3H_6$ [14], CH_4+CF_4 [15] mixed-gas hydrate systems. Although each simple hydrate constructs s-I, the crystal structure of the mixed-gas hydrates changes between s-I and s-II depending on the mole fraction of guest species (water free). The transition to s-II hydrates draws much attention to increase storage capacity of CH_4 in s-II hydrates compared with that of s-I hydrates.

In this section, I adopted $CH_4+C_2H_6$ mixed-gas as a simple model of natural gas and 1,1-DMCH as a thermodynamic promoter. I measured the four-phase (HL₁L₃G) equilibrium relation in the $CH_4+C_2H_6+1,1$ -DMCH+water system at T = 288.15 K. In addition, the hydrate structures and the cage occupancies of guest species were investigated by means of Raman spectroscopy.

III.4.3. Experimental

Materials

The materials used in the present study are summarized in **Table III-4-1**. All of them were used without further purification.

Chemical name	Source	Mole fraction purity	
CH ₄	Liquid Gas Co., Ltd.	> 0.9999	
C2H6	Takachiho Trading Co., Ltd.	> 0.999	
1,1-DMCH	Sigma-Aldrich Co. LLC.	> 0.99	
Distilled water	Wako Pure Chemical Ind., Ltd.	> 0.9999	

Table III-4-1. Information on the chemicals used in the present study.

Apparatus

The experimental apparatus for isothermal phase equilibrium measurement and Raman spectroscopy used in the present study are the same as described in **Chapter II Section 3**.

Procedures

The experimental procedures for isothermal phase equilibrium measurement and Raman spectroscopy performed in the present study are the same as described in **Chapter II Section 3**.

III.4.4. Results and Discussion

Figure III-4-1 shows Raman spectra of the C–H stretching, C–C stretching, and ringbreathing vibration modes of guest species in hydrate phase. These spectra were obtained under the four-phase (HL₁L₃G) equilibrium conditions for the CH₄+C₂H₆+1,1-DMCH+water system at T =288.15 K. The Raman peaks indicated by closed inverse triangles are derived from the intramolecular vibration mode of 1,1-DMCH in the U-cage. The single peak detected at $\Delta \nu = 751$ cm⁻¹ is derived from the sapphire window.



Figure III-4-1. Raman spectra of the C–H stretching, C–C stretching, and ring-breathing vibration modes of guest species in hydrate phase for CH₄+C₂H₆+1,1-DMCH+water system under four-phase (HL₁L₃G) equilibrium conditions at T = 288.15 K. Closed inverse triangles stand for the peaks derived from 1,1-DMCH in U-cage. The broken line stands for the contribution ($\Delta \nu = 751$ cm⁻¹) from sapphire window.

At p = 6.74 MPa and $y_{C2H6} = 0.08$, a peak derived from the ring-breathing vibration mode of 1,1-DMCH was observed at $\Delta v = 708$ cm⁻¹, which indicates that 1,1-DMCH is enclathrated in U-cages of s-H hydrates, while at $\Delta v = 705$ cm⁻¹ in the 1,1-DMCH-rich liquid phase [16]. The Raman shifts of 1,1-DMCH are consistent with the values reported in the previous study [16]. No remarkable Raman peak originated from the C–C stretching vibration mode of C₂H₆ was detected. A single peak was detected at $\Delta v = 2913$ cm⁻¹, which corresponds to the C–H stretching vibration mode of the CH₄ in the S- and S'-cages [6,16,17]. Other peaks at p = 6.74 MPa and $y_{C2H6} = 0.08$ are derived from the intramolecular vibration modes of 1,1-DMCH. The Raman shifts and spectral shapes are similar to those obtained from the CH₄+1,1-DMCH hydrate crystal [16].

In the composition range over $y_{C2H6} = 0.25$, a single peak derived from C₂H₆ got remarkable instead of 1,1-DMCH. At p = 5.82 MPa and $y_{C2H6} = 0.25$, the spectral shape of the hydrate phase is completely different from that of the s-H hydrate at p = 6.74 MPa and $y_{C2H6} = 0.08$. The Raman peak observed at $\Delta v = 992$ cm⁻¹ corresponds to that of C₂H₆ encaged in the L-cages of s-II hydrates [10,13]. The peaks detected at $\Delta v = 2903$ cm⁻¹ and 2914 cm⁻¹ are derived from CH₄ in the L- and Scages of s-II hydrates, respectively. The peaks detected at $\Delta v = 2887$ cm⁻¹ and 2942 cm⁻¹ are Fermi resonance peaks originated from C₂H₆ in L-cages. The Raman shifts are similar to those obtained from the s-II CH₄+C₂H₆ hydrate [10,13].

With further increase in y_{C2H6} , the single peak derived from the C₂H₆ molecule encaged in the M-cage of s-I hydrate [10,13,18] was detected at $\Delta \nu = 1000$ cm⁻¹ and the crystal structure of the mixed-gas hydrate transformed from s-II to s-I. Furthermore, there is no peak around $\Delta \nu = 1022$ cm⁻¹ of C₂H₆ forced to be enclathrated in the S-cage [18]. Therefore, the C₂H₆ molecule occupies only the large cages of s-I and s-II hydrates under the present experimental conditions. The CH₄ molecules are encaged in both the large (M-, L-cages) and small cages as shown in **Figure III-4-1**. The Raman peaks detected at $\Delta \nu = 2904$ cm⁻¹ and 2915 cm⁻¹ are derived from CH₄ in the M- and S-cages of s-I hydrates [6,10,11,13,17,19]. The peaks detected at $\Delta \nu = 2891$ cm⁻¹ and 2947 cm⁻¹ are the Fermi resonance peaks of C₂H₆ in the M-cages [10,11,13]. It is concluded that the crystal structure of the mixed-gas hydrates in the CH₄+C₂H₆+1,1-DMCH+water quaternary system changes from s-I to s-II and back to s-I [10–13].

The pressure-composition projection for the CH₄+C₂H₆+1,1-DMCH+water system containing gas hydrates at T = 288.15 K is shown in **Figure III-4-2**. **Table III-4-2** summarizes the relation of equilibrium pressure and composition of gas phase. The three-phase equilibrium conditions for the CH₄+C₂H₆ mixed-gas hydrate system [13] are also shown in **Figure III-4-2** to compare with those of the present system. The equilibrium pressure at $y_{C2H6} = 0$ (i.e. the s-H CH₄+1,1-DMCH hydrate) agrees well with that reported in the previous study [9].



Figure III-4-2. Four-phase (HL₁L₃G) equilibrium pressure-composition projection for the CH₄+C₂H₆+1,1-DMCH+water system and three-phase (HL₁G) equilibrium pressure-composition projection for the CH₄+C₂H₆+water system [13] at T = 288.15 K.

<i>p</i> / MPa ^{<i>a</i>}	УС2н6 ^а	<i>p</i> / MPa ^a	УС2н6 ^а	<i>p</i> / MPa ^a	УС2H6 ^{<i>a</i>}
$H_H L_1$	L ₃ G	$H_{II}L_1$	L ₃ G	$H_I L_1$	L ₃ G
5.78	0	7.30	0.10	5.59	0.29
6.25	0.04	7.16	0.11	5.29	0.31
6.27	0.05	7.02	0.12	5.24	0.31
6.49	0.06	6.89	0.13	5.07	0.33
6.71	0.08	6.41	0.16	4.87	0.34
6.72	0.09	6.19	0.18	4.82	0.35
6.94	0.10	6.17	0.19	4.74	0.37
7.01	0.11	6.11	0.20	4.58	0.38
		5.99	0.22	4.29	0.44
		5.83	0.25	4.25	0.44
		5.63	0.27	4.19	0.45
		5.78	0.27	4.14	0.47
				4.09	0.48
				3.54	0.64
				3.26	0.82

Table III-4-2. Four-phase (HL₁L₃G) equilibrium data for CH₄+C₂H₆+1,1-DMCH+water system at T = 288.15 K.

^{*a*} u(p) = 0.04 MPa, u(T) = 0.02 K, and $u(y_{C2H6}) = 0.005$.

Large pressure reduction from the CH₄+C₂H₆+water ternary system was observed in the composition range of $y_{C2H6} = 0-0.10$, which results from the formation of the s-H hydrate. The equilibrium pressure increases with a rise in the mole fraction of C₂H₆. If the C₂H₆ molecule occupies none of cages in the s-H hydrate, C₂H₆ would behave a diluent gas toward s-H CH₄+1,1-DMCH hydrate formation in the composition range of $y_{C2H6} = 0-0.10$. To evaluate cage occupancy of C₂H₆ in the s-H hydrate, the following thermodynamic prediction [20,21] based on the fugacity of CH₄ was performed.

In the CH₄+1,1-DMCH+water ternary system without C_2H_6 , the equilibrium fugacity of CH₄ in the gas phase can be evaluated at a given temperature as:

$$f_{\rm CH4}^{\rm o} = \phi_{\rm CH4}^{\rm o} p^{\rm o}$$
 (Equation III-4-1)

where f_{CH4}^{o} and ϕ_{CH4}^{o} are the fugacity and fugacity coefficient of CH₄ at four-phase (HL₁L₃G) equilibrium pressure p^{o} , respectively. Superscript o represents the absence of C₂H₆. In the present study, the fugacity coefficient was calculated by use of the Soave-Redlich-Kwong equation of state [22].

In the $CH_4+C_2H_6+1$,1-DMCH+water quaternary system, the equilibrium fugacity of CH_4 in the gas phase can be evaluated as:

$$f_{\rm CH4} = y_{\rm CH4} \phi_{\rm CH4} p \qquad (Equation III-4-2)$$

where y_{CH4} is the equilibrium composition of CH₄ in the gas phase and *p* stands for the four-phase (HL₁L₃G) equilibrium pressure (total pressure).

Supposing that the gas hydrate generated from the $CH_4+C_2H_6+1,1$ -DMCH+water quaternary system is the s-H CH₄+1,1-DMCH hydrate and the cage occupancies of CH₄ (in S- and S'-cages) and 1,1-DMCH (in U-cage) remain stationary, the fugacity of CH₄ calculated from **Equations III-4-1 and 2** should coincide with each other. In other words, the CH₄+1,1-DMCH hydrate is supposed to be generated when the fugacity of CH₄ in the gas mixture exceeds the equilibrium fugacity of CH₄+1,1-DMCH hydrate system.

Equation III-4-3 is derived from Equations III-4-1 and 2 as:

$$p = \frac{\phi_{\text{CH4}}^{\circ}}{\phi_{\text{CH4}}} \frac{p^{\circ}}{y_{\text{CH4}}}$$
(Equation III-4-3)

At a given gas phase composition, the equilibrium pressure is obtained by the trial and error method from **Equation III-4-3** on the assumption that the equilibrium fugacity of CH₄ in gas mixture is equal to that ($f_{CH4} = 5.19$ MPa) of CH₄+1,1-DMCH hydrate system in the whole composition range. The fugacity coefficient of CH₄ in the gas mixture was calculated by the Soave-Redlich-Kwong equation of state [22] with the ordinary mixing rule (k = -0.0078 for CH₄+C₂H₆) [23]. The critical constants of CH₄ and C₂H₆ were obtained from the reference [24].

Figure III-4-3 shows the comparison of experimental data with the predicted iso-fugacity curve in s-H hydrate region. The experimental pressures shift to the high-pressure side of the predicted pressures in this region. This result implies that the hydrate phase would be changed from the $CH_4+1,1$ -DMCH hydrate to $CH_4+C_2H_6+1,1$ -DMCH hydrate. In other words, there is a possibility that C_2H_6 is encaged in any cages of the s-H hydrate.



Figure III-4-3. Experimental data and predicted iso-fugacity curve in s-H hydrate region.

The four-phase equilibrium curve shows a discontinuous change in gradient at around $y_{C2H6} = 0.10$. Based on the results of Raman spectroscopic analysis, the crystal structure of the mixed-gas hydrates transforms from s-H to s-II at the composition. Beyond the composition of $y_{C2H6} = 0.10$, the equilibrium pressures are close to those for the CH₄+C₂H₆ mixed-gas hydrate system [13]. The equilibrium curve represented in **Figure III-4-2** shows a discontinuous change in gradient again at the composition of $y_{C2H6} = 0.29 \pm 0.04$. The Raman spectrum obtained at p = 5.82 MPa and $y_{C2H6} = 0.25$ indicates the formation of the s-II hydrate, while that obtained at p = 5.10 MPa and $y_{C2H6} = 0.33$ indicates the formation of the s-I hydrate. Therefore, the discontinuous point corresponds to the hydrate structural phase transition point from s-II to s-I. In the case without 1,1-DMCH (i.e. the CH₄+C₂H₆ mixed-gas hydrate system), it is reported that the transition point from s-II to s-I exists at around $y_{C2H6} = 0.40$ at T = 288.15 K [13]. The shift of transition point may be caused by the existence of 1,1-DMCH, but is not clear at this stage.

The crystal structure of the mixed-gas hydrate is s-I in the composition range from $y_{C2H6} = 0.29-1$. The equilibrium pressures are almost consistent with those of the CH₄+C₂H₆ mixed-gas hydrate system [13]. Therefore, it is suggested that the addition of 1,1-DMCH hardly affects on the equilibrium pressure of s-I CH₄+C₂H₆ mixed-gas hydrate at $y_{C2H6} > 0.29$ except liquid C₂H₆ phase appears instead of gas phase under the conditions of $y_{C2H6} = 0.82-1.0$.

III.4.5. Conclusions

Isothermal phase equilibria have been investigated for the $CH_4+C_2H_6+1,1$ -DMCH hydrate system at T = 288.15 K. In addition, crystal structures of the mixed-gas hydrates have been identified by means of Raman spectroscopy. The most important finding is that the existence of 1,1-DMCH results in a different phase behavior from the $CH_4+C_2H_6$ mixed-gas hydrate system. The thermodynamically stable phase in the $CH_4+C_2H_6+1,1$ -DMCH+water quaternary system is transformed from s-H to s-I via s-II with increase in C_2H_6 concentration, while from s-I to s-II and back to s-I without 1,1-DMCH. In the present system, CH_4 and C_2H_6 are enclathrated in the s-I and s-II hydrates. On the other hand, CH_4 and 1,1-DMCH are mainly enclathrated in the s-H hydrate. In the s-H hydrate, there is a possibility that C_2H_6 can be enclathrated in hydrate cages.

III.4.6. Nomenclatures

ν	Wavenumber [m ⁻¹]
$\phi_{ m CH4}$	Fugacity coefficient of CH ₄ [-]
fсн4	Fugacity of CH ₄ [Pa]
k	Binary parameter between CH ₄ and C ₂ H ₆ [-]
р	Pressure [Pa]
Т	Temperature [K]
и	Standard uncertainties of pressure [Pa], temperature [K], and mole fraction [-]
УСН4	Mole fraction of CH ₄ in gas phase (water and 1,1-DMCH free) [-]
УС2Н6	Mole fraction of C ₂ H ₆ in gas phase (water and 1,1-DMCH free) [-]
G	Gas phase
Н	Hydrate phase
HI	s-I hydrate phase
H_{II}	s-II hydrate phase
$H_{\rm H}$	s-H hydrate phase
L_1	Aqueous phase
L ₃	1,1-DMCH-rich liquid phase

III.4.7. References

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Chapter IV

General Conclusions

IV.1. Conclusions in the Thesis

Many kinds of mixed hydrate systems were investigated as described in **Chapters II and III**. Crystal structures of mixed hydrates were directly analyzed with PXRD. Thermodynamic stability boundaries of the mixed hydrates and cage occupancies of guest species were also clarified. In this thesis, mixed hydrates were classified into two types according to the cage occupancy of guest species; competitive cage occupancy and compartmental cage occupancy.

Competitive-Type Mixed Hydrates

Structural phase transitions were observed in the isotherms of some mixed hydrate systems, which indicates that a mixed hydrate belongs to thermodynamically stable structure at a given pressure, temperature, and composition. In some cases, a structural phase transition causes heterogeneous azeotropic-like behavior where equilibrium curve of hydrate phase changes discontinuously. Local pressure minima, observed in the isotherms of some mixed hydrate systems, are due to negative azeotropic-like behavior where equilibrium composition of gas phase coincides with that of hydrate phase. Besides, the higher the temperature is, the more remarkable the negative azeotropic-like behavior is. From the relation between temperature and equilibrium composition at the local pressure minima in s-I and s-II hydrate regions, small and large guest species in the certain mixed gas hydrate systems almost compartmentally occupy small and large cages of s-I and s-II hydrates at the local pressure minima. At this point, the mixed hydrate can be treated as if it is the simple hydrate and can form at relatively low pressure. Therefore, it is significantly valuable for utilization.

Compartmental-Type Mixed Hydrates

Thermodynamic stabilities of s-II and s-H hydrates remarkably depend on the large guest species. In this study, it was clarified that small guest species also dominates whether large guest species is enclathrated in large cage or not. CH₄ is more helpful than CO₂ for the enclathration of large guest species in s-H hydrate. It can be commonly stated that c-C₅H₁₀ derivatives work as effective thermodynamic promoters, especially c-C₅H₉F is superior. The extraordinary stability of s-H mixed hydrate at high pressures and temperatures was firstly proved. The isotherm in the quaternary mixed hydrate system, including natural gases and large guest species, showed the phase transitions among s-I, s-II, and s-H hydrates depending on the composition of gas phase. Based on the results on the s-H hydrates in this chapter, I found out the possibility that small and large guest species may competitively occupy U-cage at a certain thermodynamic condition, besides, some

small guest species multiply occupy a U-cage as shown in **Figure IV-1**. This is significantly interesting from the viewpoint of properties of s-H hydrates.

IV.2. Suggestion for the Future Works

In this thesis, the mixed hydrates composed of target gas and thermodynamic promoter were classified into two types, competitive cage occupancy and compartmental cage occupancy. Competitive and compartmental-type mixed hydrates are controlled for each application. However, the competitive cage occupancy appears in some compartmental-type mixed hydrate system and vice versa. The findings in this thesis reveal that the cage occupancy of guest species in mixed hydrates is complicated and dependent on pressure, temperature, and the combination of small and large guest species. Further investigation of the thermodynamic properties of unclarified mixed hydrate is really important for its utilization. Moreover, only experimental results on the thermodynamic properties are not sufficient at all. It is necessary to collect other properties, for instance, kinetic and rheological properties of hydrate such as formation or dissociation rates, viscosities, and cohesion. I deeply wish that, digesting the knowledge obtained from these studies, the investigation on hydrate is developed with many kinds of approach and hydrate is utilized practically in chemical and physical processes.



Figure IV-1. Schematic illustration of competitive and multiple U-cage occupancy by small guest species.

List of Publications

[1] <u>Yuuki Matsumoto</u>, Hiroshi Miyauchi, Takashi Makino, Takeshi Sugahara, Kazunari Ohgaki, "Structural Phase Transitions of Methane+Ethane Mixed-Gas Hydrate Induced by 1,1-Dimethylcyclohexane", *Chemical Engineering Science*, **66**, 2672–2676 (2011).

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List of Presentations

[1] ○<u>Yuuki Matsumoto</u>, Hiroshi Miyauchi, Takashi Makino, Takeshi Sugahara, Kazunari Ohgaki, "Structural Phase Transitions of Methane+Ethane Mixed-Gas Hydrate Induced by 1,1-Dimethylcyclohexane", *7th International Conference on Gas Hydrate*, 428.00, Edinburgh, Scotland, United Kingdom, July 17–21, 2011, Oral Presentation.

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Yuuki Matsumoto

Environmental Physical Chemistry Group, Division of Chemical Engineering, Department of Materials Engineering Science, Graduate School of Engineering Science, Osaka University