Molecular and Isotopic Composition of Volatiles in Gas Hydrates and in Sediment from the Joetsu Basin, Eastern Margin of the Japan Sea

Akihiro Hachikubo 1,*, Katsunori Yanagawa 2, Hitoshi Tomaru 3, Hailong Lu 4 and Ryo Matsumoto 5

1 Environmental and Energy Resources Research Center, Kitami Institute of Technology, 165 Koen-cho, Kitami 090-8507, Japan
2 Faculty of Social and Cultural Studies, Kyushu University, 744 Motoooka, Nishi-ku, Fukuoka 819-0395, Japan; E-Mail: kyanagawa@scs.kyushu-u.ac.jp
3 Department of Earth Sciences, Graduate School of Science, Chiba University, 1-33 Yayoi-cho, Inage-ku, Chiba 263-8522, Japan; E-Mail: tomaru@chiba-u.jp
4 Department of Energy and Resource Engineering, College of Engineering, Peking University, Beijing 100871, China; E-Mail: hlu@pku.edu.cn
5 Gas Hydrate Laboratory, Organization for the Strategic Coordination of Research and Intellectual Properties, Meiji University, 1-1 Kanda-Surugadai, Chiyoda-ku, Tokyo 101-8301, Japan; E-Mail: ryo_mat@meiji.ac.jp

* Author to whom correspondence should be addressed; E-Mail: hachi@mail.kitami-it.ac.jp; Tel.: +81-157-26-9522; Fax: +81-157-26-9534.

Academic Editor: Richard B. Coffin

Received: 31 January 2015 / Accepted: 12 May 2015 / Published: 25 May 2015

Abstract: Hydrate-bearing sediment cores were retrieved from the Joetsu Basin (off Joetsu city, Niigata Prefecture) at the eastern margin of the Japan Sea during the MD179 gas hydrates cruise onboard R/V Marion Dufresne in June 2010. We measured molecular and stable isotope compositions of volatiles bound in the gas hydrates and headspace gases obtained from sediments to clarify how the minor components of hydrocarbons affects to gas hydrate crystals. The hydrate-bound hydrocarbons at Umitaka Spur (southwestern Joetsu Basin) primarily consisted of thermogenic methane, whereas those at Joetsu Knoll (northwestern Joetsu Basin, about 15 km from Umitaka Spur) contained both thermogenic methane and a mixture of thermogenic and microbial methane. The depth concentration profiles of methane, ethane, propane, CO2, and H2S in the sediments from the Joetsu Basin...
area showed shallow sulfate–methane interface (SMI) and high microbial methane production beneath the SMI depth. Relatively high concentrations of propane and neopentane (2,2-dimethylpropane) were detected in the headspace gases of the hydrate-bearing sediment cores obtained at Umitaka Spur and Joetsu Knoll. Propane and neopentane cannot be encaged in the structure I hydrate; therefore, they were probably excluded from the hydrate crystals during the structure I formation process and thus remained in the sediment and/or released from the small amounts of structure II hydrate that can host such large gas molecules. The lower concentrations of ethane and propane in the sediment, high $\delta^{13}$C of propane and isobutane, and below-detection normal butane and normal pentane at Umitaka Spur and Joetsu Knoll suggest biodegradation in the sediment layers.

**Keywords:** gas hydrate; Japan Sea; stable isotope; neopentane; biodegradation

1. **Introduction**

Gas hydrates are crystalline clathrate compounds consisting of water and gas molecules that form at low temperatures and high pressures [1]. Natural gas hydrates are found worldwide in continental margin sediments [2–4] and in near-surface sediments associated with active gas plume that vent from the seafloor [5–7]. Natural gas hydrates are considered potential energy resources and are large reservoirs of methane (C1), and their dissociation may cause submarine geohazards and contribute to global warming [8–10].

Crystallographic structures of natural gas hydrate are usually either structure I (sI), which is composed of two 12-hedra and six 14-hedra with space group Pm3n, or structure II (sII), which is composed of sixteen 12-hedra and eight 16-hedra with space group Fd3m [1]. C1 and ethane (C2) are both known to form sI hydrates; however, certain compositions of C1 and C2 will form sII hydrates [11,12], whereas propane (C3) and isobutane (i-C4) is incorporated only in sII. The concentrations of C2 and C3 in hydrate-bound gas from the Gulf of Mexico represent 3%–5% and >15% of the total gas, respectively [13]. Gas hydrate from the Sea of Marmara contained high concentrations of C3 (18.8%) and i-C4 (9.5%) [14]. Mixed-gas (C1 and C2) hydrates in Lake Baikal belonged to the sII hydrate and contained 0.026%–0.064% of neopentane (neo-C5, 2,2-dimethylpropane) [15,16], which can be encaged in the large cages of sII [17].

Shallow gas hydrates were recently found in the Joetsu Basin at the eastern margin of the Japan Sea, where gas venting was observed on echo-sounder images [18,19]. Gas hydrate was recovered from the sea floor [20], and $\delta^{13}$C of hydrate-bound C1 was from ~37.3‰ to ~37.1‰ [20], suggesting thermogenic origin in the criteria of the Bernard plot [21]. However, C2 and C3 concentrations of hydrate-bound gas at Umitaka Spur and Joetsu Knoll (Figure 1) were very low; the molar ratio C1/(C2 + C3) ranged from 1000 to 10,000 [22]. The Sado Nansei Oki drillings at Umitaka Spur conducted by the Ministry of Economy, Trade and Industry, Japan revealed C1/(C2 + C3) values less than 100 at depths 1143–2016 m below the sea floor [20], indicating either that C2 and C3 concentrations are reduced in the ascending fluid during migration or that the addition of microbial C1 is significant in higher strata. The depth profiles of sediment gases at Umitaka Spur and Joetsu Knoll were reported in our previous
work [23]; however, molecular and isotopic compositions of hydrate-bound gas were not reported yet and the effect of low concentration of higher hydrocarbons on gas hydrates has not been discussed. In this study, we investigate the molecular and stable isotope compositions of hydrate-bound gas and gas in sediment cores (headspace gas) retrieved from Umitaka Spur and Joetsu Knoll and focus on C₂, C₃, and higher hydrocarbons, those may change crystallographic structure of the shallow gas hydrates in these areas.

**Figure 1.** Locations of the coring sites in the Joetsu Basin area, Japan Sea. The sites of hydrate-bearing cores are highlighted.
2. Study Sites and Sediment Cores

Umitaka Spur and Joetsu Knoll are located in the Joetsu Basin off the city of Joetsu (Niigata Prefecture, Figure 1). High P-wave velocities at the mounds and pockmarks on the Umitaka Spur seafloor were reported, suggesting the existence of gas hydrate [24]. Echo sounding of the sea bottom at Umitaka Spur and Joetsu Knoll showed large-scale gas venting from the sea floor [18]. Depth profiles of sulfate in the pore water of the sediment cores indicated shallow (<5 m) sulfate–methane interface (SMI; where sulfate and methane are both consumed to depletion at the base of the sulfate reduction zone) in the Joetsu Basin [25], indicating high flux of hydrocarbons and other organic matter. High heat flow was observed in a restricted area on the sea floor around the gas seep sites at Umitaka Spur (maximum value of 1590 mW m$^{-2}$) and Joetsu Knoll (maximum value of 519 mW m$^{-2}$) compared to the average value obtained on a normal muddy seafloor in these areas (98 ± 13 mW m$^{-2}$), indicating local fluid migration from the deep subsurface layer [26].

Previous study considered origin of hydrate-bound C$_1$ at Umitaka Spur and Joetsu Knoll as thermogenic one (C$_1$ $\delta^{13}$C ranged from −39.0 to −34.9‰ V-PDB) and mixed gas of microbial and thermogenic ones (C$_1$ $\delta^{13}$C ranged from −57.9 to −51.8‰ V-PDB), respectively [22]. Because the content of total organic carbon (TOC) ranged from 0.58% to 1.55% in the shallow sediment layer (0–9 m depth, [27]) and from 0.90% to 2.98% in the deep layer (1360–2088 m depth, mixed Type II/III kerogen, [20]), the potential of C$_1$ generation via microbial and thermogenic processes is high.

Sediment cores were retrieved from Umitaka Spur, Joetsu Knoll, Joetsu Basin and its peripheral area (Figure 1) during the MD179 cruise onboard R/V Marion Dufresne in June 2010. A Calypso giant piston corer enabled us to obtain sediment cores up to 40 m in length. A Calypso square box corer (CASQ, 25 cm × 25 cm × 12 m) was also used. Out of the 20 sediment cores obtained, four contained hydrates: MD179-3305G and MD179-3306 (located 300 m apart) from Umitaka Spur, and MD179-3317 and MD179-3318C (located 5 km apart) from Joetsu Knoll. Hydrates were virtually absent in the other sediment cores. MD179-3305G was retrieved by a gravity corer, and a few samples of the gas hydrates were taken for gas analysis. All the hydrate-bearing cores except MD179-3318C lost several meters of the top sediment layers because the lead weight section was disconnected from the core barrel due to a technical problem; the gas hydrates dissociated in the corer and the core top was displaced during the onboard recovery process. The top depths of the gas hydrates were estimated to be 4.5 mbsf (MD179-3306), 31.5 mbsf (MD179-3317), and 1.5 mbsf (MD179-3318C). The depth of gas hydrate in MD179-3305G was unknown.

3. Sampling Methods and Analysis

3.1. Onboard Gas Sampling

The gas sampling procedures for the hydrate-bound and headspace gases were the same as those used in our previous studies [28,29]. The hydrate-bound gases were collected using a 50-mL plastic syringe and stored in 5-mL vials sealed with butyl septum stoppers. Several samples were taken from each hydrate-bearing sediment core. First, we placed small pieces of hydrate sample into a 50-mL plastic syringe, pushed the cylinder to reduce the dead volume, and connected the syringe to a 5-mL vial with a needle. Another needle was connected to the vial to flush the air inside. Each vial was thus filled with
hydrate-bound gas without sediment particles or water. Consequently, this method contributed to prevention of microbial activity during storage. The gas dissociated from hydrate flushed air in the system, and the concentration of air in the vial was less than 2%.

Headspace gas method was employed to know the depth profiles of each gas component in the sediment cores. 10 mL of sediment, 9.5 mL of saturated aqueous solution of NaCl, and 0.5 mL of preservative (10 wt % aqueous solution of benzalkonium chloride, [30]) were introduced into 25-mL vials to create a 5-mL headspace. The headspace was flushed with helium, the carrier gas used in the gas chromatography, to reduce air contamination. Although the flushing process may reduce gas concentrations from their native values, differences in concentration and stable isotopes ($\delta^{13}C$ and $\delta^{2}D$) of C1 between flushed and non-flushed samples were undetectable [31]. Hence, we can discuss the trend of their depth profiles using headspace gas data. The vials were shaken well and stored upside down until they were analyzed.

In some of the gas-rich cores (MD179-3296, 3299, 3301, 3304, 3308, 3313, and 3317, see Figure 1), voids formed in the plastic core liner during their retrieval from the core barrel. The gas in the voids was collected by modifying the method of [32]. The surface temperature of the plastic core liner was measured with an infrared thermal imaging camera to locate the voids. Small holes several meters apart were then drilled into the liner. Gas samples collected through these holes with a 50-mL plastic syringe were stored in evacuated 5-mL glass vials. Because the concentration of hydrocarbons in the void gas was considerably higher than that obtained by the headspace gas method, the void-gas method is suitable for stable carbon and hydrogen isotope analysis on non-C1 hydrocarbons, which are typically present in low concentration. Isotopic difference between void and headspace gases was within 2‰ for C1 $\delta^{13}C$ and 3‰ for C1 $\delta^{2}D$ [23]. In this study, we obtained the stable carbon isotope signature of C2 by the void-gas method.

### 3.2. Analytical Methods

The molecular composition of the gas samples was determined using a gas chromatograph (GC-14B, Shimadzu, Kyoto, Japan) equipped with a packed column (Shimadzu Sunpak-S; 2 m length, 3 mm ID), a thermal conductivity detector (TCD) for detecting CO2, H2S, and high concentrations (>0.1 mol% of the sample gas) of C1, and a flame ionization detector (FID) for detecting low concentrations (<0.1 mol% of the sample gas) of hydrocarbons (C1–C5). The TCD and FID were connected in series. The detection limit was 0.00005 mol% (C1–C3) and 0.0005 mol% (C4–C5). The analytical error estimated by multiple injections of standard gases was less than 1.2% for each gas component.

Stable carbon and hydrogen isotopic ratios of the hydrocarbons and CO2 were measured using a continuous-flow isotope-ratio mass spectrometer (CF-IRMS, DELTAplus XP, Thermo Fisher Scientific, Waltham, MA, USA) coupled with a gas chromatograph (Trace GC, Thermo Fisher Scientific) via a combustion/pyrolysis reactor (GC-C/TC III, Thermo Fisher Scientific). The gas chromatograph was equipped with a Carboxen-1006 PLOT capillary column (30 m length, 0.32 mm ID, 15 μm film thickness, Sigma-Aldrich, St. Louis, MO, USA). In the case of samples with low C1 concentration, a Sigma-Aldrich Carboxen-1010 PLOT capillary column (30 m length, 0.32 mm ID, 15 μm film thickness) was also used to separate air components from hydrocarbons. A PoraPLOT Q capillary column
(27.5 m length, 0.32 mm ID, 10 μm film thickness, Agilent Technologies, Santa Clara, CA, USA) was used for higher hydrocarbons (C4–C8). Stable isotope compositions are reported as δ values (‰):

\[
\delta_{\text{sample}} = \left( \frac{R_{\text{sample}} - R_{\text{standard}}}{R_{\text{standard}}} \right) \times 1000 \text{ (‰)}
\]

where \( R \) denotes the \(^{13}\text{C}/^{12}\text{C} \) or D/H ratio. \( \delta^{13}\text{C} \) and \( \delta D \) are given with reference to the V-PDB and V-SMOW standards, respectively, determined by using NIST RM8544 (NBS19) for \( \delta^{13}\text{C} \) and NIST RM8561 (NGS3) for \( \delta D \). The analytical precision was 0.3‰ for \( \delta^{13}\text{C} \) of C1–C3, 1‰ for \( \delta^{13}\text{C} \) of C4–C8, and 2‰ for \( \delta D \).

4. Results

Figure 2a–c show the depth profiles of the concentration (C1, C2, C3, CO2, and H2S) of headspace gases, C1/(C2 + C3) values calculated from the concentration of C1, C2, and C3, and stable carbon and hydrogen isotopes of C1, C2, and CO2. The concentration of C1 drastically increased with depth toward the SMI, with the peak concentration appearing at around 5–8 mbsf and then decreasing slightly. The C1 concentration profiles indicate that the SMI is around 0.5–5 mbsf in the Joetsu Basin area including Umitaka Spur and Joetsu Knoll. The trend of C2 and C3 concentration profiles was similar to that of C1; however, their concentrations below the SMI differed between the sediment cores. C1/(C2 + C3) values showed a peak near 5–8 m, which agrees with the peak of the C1 concentrations. C1/(C2 + C3) values below the SMI were less than 10,000 in the areas of Umitaka Spur (Figure 2a) and Joetsu Knoll (Figure 2b) except the cores MD179-3320G and 3324C, and more than 10,000 in the other cores of Joetsu Basin (Figure 2c). The CO2 concentrations of the all cores simply increased with depth, and the H2S concentrations showed peaks at their SMI depths.

The trend of C1 \( \delta^{13}\text{C} \) profiles were similar to those of CO2 \( \delta^{13}\text{C} \), increasing with depth below the SMI. Negative C1 \( \delta^{13}\text{C} \) and CO2 \( \delta^{13}\text{C} \) peaks were observed around the SMI, which agreed with the high H2S concentrations. High C1 \( \delta^{13}\text{C} \) (more than −45‰ V-PDB) was observed in the hydrate-bearing cores MD179-3306 and 3318C. C1 \( \delta^{13}\text{C} \) at the depth of 10 mbsf ranged from −66.7‰ to −57.8‰ at Umitaka Spur except the core MD-179-3306, from −74.5‰ to −58.3‰ at Joetsu Knoll, and from −83.5‰ to −77.4‰ at the other sites in Joetsu Basin. On the other hand, C1 \( \delta D \) simply decreased with depth except the core MD179-3306. C1 \( \delta D \) for MD179-3306 was distinctly large and ranged from −165.9‰ to −157.8‰, while those for other cores ranged from −207.4‰ to −175.8‰. C1 \( \delta^{13}\text{C} \), C1 \( \delta D \), and C2 \( \delta^{13}\text{C} \) of hydrate-bound and headspace gases were almost the same with each other. On the contrary, distinct differences were found in CO2 \( \delta^{13}\text{C} \); those of hydrate-bound gas was 10‰ lower (MD179-3306), 8‰ lower (MD179-3317), and 27‰ higher (MD179-3318C) than those of headspace gas around the same depth.

The concentration of C2 in the hydrate-bearing cores (MD179-3306, 3317, and 3318C) below the SMI was nearly one order of magnitude higher than that in the nonhydrate-bearing cores. MD179-3297C, 3300C, 3304, and 3307C were also rich in C2. The concentrations of C3 in the three hydrate-bearing cores (MD179-3306, 3317, and 3318C) and in MD179-3304 were higher than in the nonhydrate-bearing cores. The C2-rich cores (MD179-3304 and 3317) showed \(^{13}\text{C} \) enrichment in C2, although C3 \( \delta^{13}\text{C} \) is not available. C2 \( \delta^{13}\text{C} \) of nonhydrate-bearing cores in Joetsu Basin was low, ranging from −62.8‰ to −52.6‰.
Figure 2. Cont.
Figure 2. Depth profiles of $C_1$, $C_2$, $C_3$, $CO_2$, and $H_2S$ concentrations, $C_1/(C_2 + C_3)$ values, $C_1 \delta^{13}C$, $C_1 \delta^D$, $C_2 \delta^{13}C$, and $CO_2 \delta^{13}C$ in the headspace gas. (a) Umitaka Spur area; (b) Joetsu Knoll area; (c) Joetsu Basin and its peripheral area. These depth profiles of the headspace gases were already reported [23]. The $C_1/(C_2 + C_3)$ values and stable carbon ($C_1$, $C_2$, and $CO_2$) and hydrogen ($C_1$) isotopic ratios in the hydrate-bound gas of the cores MD179-3306, 3317 and 3318C are plotted. The data of $C_2 \delta^{13}C$ was obtained by the void-gas method and other data by the headspace gas method. The depth data of $C_2 \delta^{13}C$ profile for MD179-3296 is unknown. The headspace gas data of MD179-3305G is missing due to sediment loss.

Molecular compositions and stable isotope signatures of hydrate-bound gases at Umitaka Spur and Joetsu Knoll are summarized in Table 1. In the entire study sites $C_1$ was the main component of the hydrate-bound gas, comprising more than 98 mol% of the total volume; the concentrations of $C_2$, $C_3$, $CO_2$, and $H_2S$ were 0.0148–0.0456 mol% ($n = 17$), 0.0001–0.0025 mol% ($n = 17$), 0.03–1.26 mol% ($n = 16$), and 0.03–0.77 mol% ($n = 8$), respectively. $C_1 \delta^{13}C$ has a wide range (−57.1‰ to −43.9‰) at Joetsu Knoll (MD179-3317 and 3318C) and a narrow range (−37.3‰ to −34.6‰) at Umitaka Spur (MD179-3305G and 3306), whereas $C_1/(C_2 + C_3)$ values concentrate from 2200 to 9700 at the both areas. $C_2 \delta^{13}C$ ranged from −31.9‰ to −21.3‰ at Joetsu Knoll and from −18.4‰ to −17.9‰ at Umitaka Spur, respectively.
Table 1. Molecular compositions (mol% of the total components) and stable carbon (C$_1$–3 and CO$_2$) and hydrogen (C$_1$) isotopic ratios of hydrate-bound gases retrieved from Umitaka Spur (MD179-3305G and 3306) and Joetsu Knoll (MD179-3317 and 3318C), Japan Sea. Because H$_2$S is highly corrosive and reaction with sampling and analytical tools was apparently not prevented, a higher portion of H$_2$S is lost during core retrieval and gas analysis compared to hydrocarbons and CO$_2$.

| Core No. Depth [mbsf] | Molecular composition | Isotopic composition |
|------------------------|------------------------|----------------------|
|                        | C$_1$ [mol%] | C$_2$ [mol%] | C$_3$ [mol%] | CO$_2$ [mol%] | H$_2$S [mol%] | C$_1$/(C$_2$+C$_3$) | C$_1$$\delta^{13}$C [%V-PDB] | C$_2$$\delta^{13}$C [%V-PDB] | C$_3$$\delta^{13}$C [%V-PDB] | CO$_2$$\delta^{13}$C [%V-PDB] | C$_1$$\delta^D$ [%V-SMOW] |
| MD179-3305G unknown depth | 98.1 | 0.0292 | 0.0019 | 1.10 | 0.77 | 3158 | −36.6 | −17.9 | 5.8 | 14.3 | −167 |
| MD179-3306 4.5 [mbsf] | 99.1 | 0.0292 | 0.0020 | 1.18 | 0.53 | 3389 | −36.0 | −18.2 | 5.7 | 18.0 | −164 |
| MD179-3317 31.5 [mbsf] | 99.3 | 0.0205 | 0.0017 | 1.26 | 0.53 | 3086 | −36.0 | −18.2 | 5.9 | 19.5 | −164 |
| MD179-3318C 1.5 [mbsf] | 99.8 | 0.0331 | 0.0018 | 0.20 | n.d. | 2859 | −34.6 | 17.3 | n.d. | 167 |

Notes: mbsf: meters below sea floor; blank: not measured; n.d.: not detected.
The molecular and stable carbon and hydrogen isotopic composition of volatiles in the headspace gas of MD179-3306 (Umitaka Spur) and 3318C (Joetsu Knoll) are summarized in Table 2. A high concentration of a compound (0.1–0.2 mol%) putatively assigned as neo-C₅ based on its relative retention time during gas chromatography was also detected. Although normal butane (n-C₄), i-C₄, isopentane (i-C₅), and normal pentane (n-C₅) were below the detection limit of the gas chromatograph, CF-IRMS (detection limit: 0.00006 mol%) detected only i-C₄ in the samples. δ¹³C of C₅–C₈ in MD179-3306 and 3318C was within −23‰ ± 5‰; however, C₃ δ¹³C and i-C₄ δ¹³C were exceptionally high (+1.1‰ to +8.8‰) compared with δ¹³C of the other hydrocarbons.

Table 2. Molecular and isotopic compositions of headspace gas of the hydrate-bearing cores (MD179-3306 and MD179-3318C).

| Core | MD179-3306 | MD179-3318C |
|------|------------|-------------|
| Depth [mbsf] | 4.50 | 5.00 | 5.95 | 7.00 | 1.50 |
| Molecular composition [mol%] | | | | | |
| C₁ | 40.7 | 42.3 | 41.4 | 28.4 | 86.9 |
| C₂ | 0.0211 | 0.0264 | 0.0255 | 0.0153 | 0.0289 |
| C₃ | 0.0037 | 0.0039 | 0.0037 | 0.0022 | 0.0063 |
| neo-C₅ | 0.1946 | 0.1298 | 0.1862 | 0.0830 | 0.0760 |
| CO₂ | 59.1 | 57.5 | 58.4 | 71.5 | 6.6 |
| H₂S | n.d. | n.d. | n.d. | n.d. | 6.3 |
| C₁/(C₂+C₃) | 1640 | 1393 | 1422 | 1624 | 2469 |
| Isotopic composition [δ¹³C ‰ V-PDB] | | | | | |
| C₁ | −33.9 | −34.6 | −34.4 | −34.3 | −45.3 |
| C₂ | −47.3 | −32.9 | −21.5 | −21.1 | −24.8 |
| C₃ | 4.5 | 4.5 | 3.9 | 2.9 | 8.8 |
| i-C₄ | 2.6 | 4.4 | 2.3 | 1.1 | |
| neo-C₅ | −23.0 | −23.3 | −22.8 | −23.2 | −21.4 |
| 2,2DMB | −18.3 | −19.0 | −19.3 | −18.6 | −15.7 |
| 2,3DMB | −21.9 | −20.5 | −21.6 | |
| n-C₆ | −23.0 | −22.6 | −23.0 | |
| n-C₇ | −28.1 | −26.6 | −27.4 | |
| n-C₈ | −26.4 | | | |
| CO₂ | 31.4 | 31.5 | 31.8 | 31.9 | −10.2 |
| Isotopic composition [δD ‰ V-SMOW] | | | | | |
| C₁ | −162 | −166 | −158 | −161 | −183 |
| neo-C₅ | −125 | −124 | −133 | −125 | −119 |

Notes: mbsf: meters below sea floor; blank: not measured; n.d.: not detected; DMB: dimethylbutane.

5. Discussion

5.1. Depth Profiles of Headspace Gas

Shallow SMI was observed in the Joetsu Basin area including Umitaka Spur and Joetsu Knoll, suggesting that the hydrocarbon flux from great depth was primarily high. The peak of the C₁ concentrations and high C₁/(C₂ + C₃) values appeared around 5–8 mbsf, indicating high microbial C₁ production (methanogenesis) just beneath the SMI. Negative C₁ δ¹³C and CO₂ δ¹³C peaks and high H₂S
concentrations were observed around the SMI (Figure 2a–c), which typically results from the anaerobic oxidation of methane (AOM). C1 δ13C decreased around the SMI can be explained as a result of carbon recycling between the AOM and methanogenesis [33]. ANME-1 and ANME-2 groups were distinguished at Umitaka Spur and Joetsu Knoll [34] using the lipid biomarker signatures of the AOM communities [35]. In the same study area, the occurrence of AOM in the SMI zone is also supported by environmental DNA analysis [36]. Similar relations between the depth profiles of C1 δ13C and CO2 δ13C were also reported in the gassy sediment of Eckernförde Bay in the western Baltic Sea [37] and in the C1-rich sediment near a gas chimney in northern Gulf of Mexico [38]. Although high C1 δ13C was observed in the hydrate-bearing core MD179-3318C that indicated thermogenic origin according to empirical classifications [21,39], depleted C1/(C2+C3), C1 δ13C, and CO2 δ13C around SMI were observed even in the MD179-3318C, suggesting that C1 is basically thermogenic and the top layer around SMI depth is affected by microbial alternation.

The upward increase in the profiles of C1 δ13C and CO2 δ13C above the SMI is due to C1 oxidation near the sea floor [39]. On the other hand, the increase in the C1 δ13C below the SMI is explained by the concurrent increase in the CO2 δ13C with depth, resulting from the reduction of CO2 in the methanogenic zone and the Rayleigh process. Except for MD179-3306, the profiles of C1 δD were almost identical and gradually decreased with depth. This trend agrees fairly well with previously published δD profiles of pore water (ambient H2O) in the same cores [40]; this is because C1 δD is primarily determined by δD of H2O and H2 in the ambient water in the case of microbial C1 generation via CO2 reduction [41]. In contrast, the profile of C1 δD for MD179-3306 was distinctly larger than those of other cores, indicating that C1 is of thermogenic origin.

The concentration of C2 in the three hydrate-bearing cores below the SMI was nearly one order of magnitude higher than that in the nonhydrate-bearing cores (Figure 2a–c), and the other cores at Umitaka Spur were also rich in C2. These C2-rich cores showed 13C enrichment in C2. The concentrations of C3 in the hydrate-bearing cores and in MD179-3304 were also higher than in the nonhydrate-bearing cores. The results for C2 and C3 suggest two possibilities: (1) injections of thermogenic hydrocarbons from greater depth and (2) dissociation of tiny amounts of C2- and C3- rich sII hydrates during recovery process of sediment cores.

The concentration of C3 in MD179-3317 was conspicuously higher (0.039–0.331 μM) than in the other cores, whereas the hydrate-bound C3 in MD179-3317 was negligible (0.0001% of the hydrate-bound gas, Table 1). Because sI gas hydrates cannot encage C3, it is reasonable to assume that C3 was excluded from the hydrate formation. Similar molecular fractionations were described in previous studies [42–44].

5.2. Effect of Gas Characteristics on Hydrate Crystal

The relation between the molecular ratio of hydrate-bound hydrocarbons and the stable carbon and hydrogen isotopic ratio of individual hydrate-bound volatiles is shown in Figure 3. The Bernard plot (Figure 3a), which compares C1 δ13C with C1/(C2 + C3) [21], is useful for understanding the gas origin and its pathway. The data from stations MD179-3305G, 3306, and 3318C plot in the mixed gas field, whereas those from MD179-3317 plot near and within the field of microbial hydrocarbons (Figure 3a). These relation between C1 δ13C and C1/(C2 + C3) at Umitaka Spur and Joetsu Knoll agree well with those obtained in a previous study [22].
Figure 3. Relationship between molecular and isotopic composition of hydrate-bound hydrocarbons. (a) “Bernard plot” showing relationship between C$_1$ $\delta^{13}$C and C$_1$/(C$_2$ + C$_3$) values [21]. The data agree well with those obtained at the same area from 2005 to 2008 [22]. The headspace gas data obtained from deeper sediment layer of Umitaka Spur at depths 993–2016 m below the sea floor [20] are plotted; (b) Relationship between C$_1$ $\delta^{13}$C and C$_2$ $\delta^{13}$C, based on the classification of Milkov [4].

Figure 3b shows the relation between C$_1$ $\delta^{13}$C and C$_2$ $\delta^{13}$C. The boundaries between thermogenic and microbial origins are based on literature [4,45,46]. C$_2$ $\delta^{13}$C ranged from $-31.9\%_{o}$ to $-17.9\%_{o}$, plotted in the field of thermogenic C$_2$ [4,45,46]. Because microbial C$_2$ is generally a minor component of microbial gas (<0.1%, see Figure 3a) and is depleted in $^{13}$C [46], mixing of microbial and thermogenic hydrocarbons slightly decreases the C$_2$ $\delta^{13}$C and strongly decreases the C$_1$ $\delta^{13}$C.

The molecular ratios of hydrate-bound hydrocarbons and stable carbon isotope signatures of C$_1$ shown in Figure 3a indicate a mixed gas origin for the hydrate at Umitaka Spur, implying depletion of C$_2$ and C$_3$ from the thermogenic gas field in Figure 3a (C$_1$/(C$_2$ + C$_3$) < 100). C$_1$/(C$_2$ + C$_3$) values were
less than 100 at depths 1143–2016 m below the sea floor at Umitaka Spur [20]. Although thermogenic gas might contain significant amounts of C\textsubscript{2} and C\textsubscript{3} (sometimes >10 mol%), as found in samples from the Gulf of Mexico [13,47], the northern Cascadia margin offshore Vancouver Island [48], and the Caspian Sea [6], the composition of C\textsubscript{2} and C\textsubscript{3} of hydrate-bound hydrocarbons at Umitaka Spur was less than 0.04% and 0.002%, respectively (Table 1). These results can be explained by molecular hydrocarbon fractionation during upward gas migration [49]. Several similar theories have been proposed, e.g., formation of C\textsubscript{2}-rich gas hydrates in the deep layers and C\textsubscript{2} depletion in the residual migrating gas [22], and adsorption of C\textsubscript{2} and C\textsubscript{3} to the mineral matrix during migration [30]. Assuming steady-state upward migration of thermogenic hydrocarbons, we have to consider the consumption and/or degradation of C\textsubscript{2} and C\textsubscript{3} in the deeper sediment layers. Non-C\textsubscript{1} hydrocarbon degradation [43,50,51] likely selectively decreases C\textsubscript{2} and C\textsubscript{3}.

Stable isotope signatures (C\textsubscript{1} $\delta^{13}$C, C\textsubscript{2} $\delta$D, and C\textsubscript{2} $\delta^{13}$C) of hydrate-bound and headspace gases were almost the same with each other (Figure 2a,b); however, distinct differences were found in CO\textsubscript{2} $\delta^{13}$C. Previous studies showed no or rather small differences in CO\textsubscript{2} $\delta^{13}$C between stable isotopes of hydrate-bound and headspace gases [42,52]. It remains unsettled and further studies are needed to find the reason why such large differences in CO\textsubscript{2} $\delta^{13}$C existed.

Although C\textsubscript{3} composition of hydrate-bound gas was very small (less than 0.0025 mol%, Table 1), we need to discuss the existence of C\textsubscript{3} in the hydrate-bound gas. C\textsubscript{1}, C\textsubscript{2}, CO\textsubscript{2}, and H\textsubscript{2}S can be encaged in the sI gas hydrate [1], but C\textsubscript{3} is exclusively found in the sII gas hydrate because of its large molecular size. It is possible that C\textsubscript{3} in these samples was a contaminant from the sediments attached to the hydrate during the preparation of the hydrate-bound gas. Based on the gas compositions, the gas hydrates investigated in this study were assumed to be mainly sI. This assumption is supported by the powder X-ray diffraction (PXRD) data for gas hydrates collected from Umitaka Spur and Joetsu Knoll in the past [53,54]. Nevertheless, we cannot exclude the potential coexistence of sI (C\textsubscript{1}-rich) and sII (C\textsubscript{3} encaged in the large cages) gas hydrates, because PXRD cannot detect small amounts of sII. As encaged C\textsubscript{3} molecules in C\textsubscript{1}-rich natural gas hydrate retrieved at the Mackenzie Delta (Onshore, Canada) were detected by Raman spectroscopy [55], heterogeneity of hydrocarbons in the different crystallographic structures is an attractive research target in the future from the viewpoint of crystal growth process of natural gas hydrates.

5.3. Neopentane and Non-Methane Hydrocarbons in the Headspace Gases

The presence of gem-dimethyl hydrocarbons was previously reported in the Deep Sea Drilling Project Holes 381 and 397 [56,57]. neo-C\textsubscript{5} is considered to form from the decomposition of gem-dimethylcycloalkanes derived from the terpenes of terrestrial organic matter [56]. Unusually high concentration of neo-C\textsubscript{5} in gases extracted from the Athabasca oil sands and proposed that neo-C\textsubscript{5} forms from the microbial degradation of bitumen [58]. Although the origin of neo-C\textsubscript{5} remains unknown, the potential involvement of microbes has been suggested [57]. Schaefer and Leythaeuser [59] explained that enrichment of neo-C\textsubscript{5} is caused by preferential diffusion due to the nearly spherical molecules and its diffusion coefficient, which is higher than that of less branched isomers.

Although neo-C\textsubscript{5} can be encaged in the large cages of sII gas hydrates [17], only trace levels of neo-C\textsubscript{5} were detected in the samples of hydrate-bound gas, perhaps because of contamination with the
sediment gas at their retrieval process. Similar to C₃, during the formation of sI gas hydrates, neo-C₅ is excluded and remains in sediment. However, negligible amounts of sII gas hydrates containing neo-C₅ could also be present, which dissociated during core retrieval. While C₃ and neo-C₅ in C₁-rich hydrocarbons decrease the equilibrium pressures and stabilize their hydrate phase, the dissociation behavior of the gas hydrate depends on the guest molecules. C₁ hydrate shows self-preservation phenomena [60] in the dissociation process; however, the hydrates with lower decomposition pressures (C₂ and C₃) do not show it [61]. Further investigations are needed to understand the behavior of the small amount of sII formers, C₃ and neo-C₅.

C₃ δ¹³C and i-C₄ δ¹³C were exceptionally high compared with δ¹³C of the other hydrocarbons as shown in Table 2. Possibly, ¹³C-depleted C₃ and i-C₄ were preferentially consumed as microbial substrates; hence, the residual C₃ and i-C₄ were enriched in ¹³C [30]. The concentration of n-C₄ was below the detection limit (0.00006 mol% of the original headspace sample) of the CF-IRMS; n-C₄ is easily transformed by hydrocarbon degraders, whereas C₂ and i-C₄ appear less affected or remain unaltered [42,50,51,62]. Therefore, it is reasonable to assume that n-C₄ was consumed primarily by biodegradation because it is more affected by biodegradation than i-C₄. δ¹³C of C₁–C₈ show a linear trend, indicating no microbial contribution in C₁. Moreover, the hydrate-bound C₁ comprised pure thermogenic gas.

![Figure 4. δ¹³C of hydrocarbons plotted in the “Natural gas plot” adapted from [63], where n is the number of the carbon atoms of individual hydrocarbon molecules. δ¹³C of C₁, C₂, C₃, n-C₄, and n-C₅ were plotted in the original natural gas plot. However, we plotted the data of i-C₄ (instead of n-C₄), neo-C₅ (instead of n-C₅), and C₆–₈ shown in Table 2. The concentrations of n-C₄, and n-C₅ were below the detection limit. Thick shaded lines show linear relation between C₂ and C₃ for MD179-3305G and 3306 (Umitaka Spur) and for MD179-3318C (Joetsu Knoll), except ¹³C-rich C₃ and i-C₄ due to potential biodegradation. C₂ δ¹³C of MD179-3306 has a large error bar because of wide distribution along with depth (Table 2).](image-url)

In contrast, these gases mixed with microbial C₁ during migration to shallow sediment layers in the Joetsu Knoll area, as C₁ δ¹³C decreased from the anticipated thermogenic C₁ value for MD179-3318C.
6. Conclusions

Molecular and stable carbon and hydrogen isotopic compositions of hydrate-bound and pore-water gas were reported for samples retrieved from Umitaka Spur and Joetsu Knoll at the eastern margin of the Japan Sea. According to empirical classifications of gas data, the hydrate-bound gas from Umitaka Spur was of thermogenic origin and that from Joetsu Knoll partly contained microbial gas. C1 concentration depth profiles showed shallow SMI in these areas, indicating high C1 flux. The molecular composition of C2 and C3 for the headspace gas of hydrate-bound cores were around a thousandth smaller than those of hydrate-bound thermogenic gases across the world, and C3 δ13C and i-C4 δ13C were exceptionally high (+1.1‰ to +8.8‰) in the pore water, suggesting that biodegradation affects these hydrocarbons in hydrate-bearing sediment systems.

High concentration of neo-C5 was detected in the pore water gases in MD179-3306 (Umitaka Spur) and 3318C (Joetsu Knoll), which suggests that neo-C5 was excluded from the hydrate crystal during the formation process and thus remained in the pore water, because neo-C5 cannot be encaged by sI gas hydrates. This is supported by the observation that the concentration of C3 in the hydrate-bearing core was higher than that in the nonhydrate-bearing cores. However, small amounts of sII gas hydrate can possibly encage these heavier hydrocarbons. Because neo-C5 forms from the microbial degradation of organic matter and is easy to diffuse in sediment layers, the accumulation of neo-C5 in subsurface sediment can affect and change the crystallographic properties of gas hydrates in marine sediments.

Acknowledgments

We appreciate the support of the crew onboard R/V Marion Dufresne during the MD179/Japan Sea Gas Hydrate cruise. We express our gratitude to the scientists onboard the MD179/Japan Sea Gas Hydrate cruise. We also thank Thomas Pape (University of Bremen, Germany) for valuable suggestions and comments. This study was mainly supported by the MH21 Research Consortium for Methane Hydrate Resources in Japan, and the analytical system was supported by the Grant-in-Aid for Scientific Research (C) 22540485 and (B) 26303021 of the Japan Society for the Promotion of Science (JSPS).

Author Contributions

Akihiro Hachikubo designed the study, performed gas analysis, and drafted the manuscript. Katsunori Yanagawa (microbiology), Hitoshi Tomaru (pore water geochemistry), and Hailong Lu (crystallography) helped to draft the manuscript from their professional point of view. Ryo Matsumoto designed the framework of the MD179 cruise. All authors discussed the results and approved the final manuscript.

Conflicts of Interest

The authors declare no conflict of interest.
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