Sources of seasonal wetland methane emissions in permafrost regions of the Qinghai-Tibet Plateau

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In this study, systematic soil methane cycle geochemical monitoring was carried out in a typical gas hydrate region in the Qinghai-Tibet Plateau. Soil gas samples were collected for hydrocarbon components and carbon isotope analysis. Meanwhile, soil-methane fluxes from the upper active layer (20–30 cm) were monitored during six months of one year. The results of this research provide evidence of a new source of methane emission from wetland soils in permafrost regions: gas hydrate release. Sites with large methane emissions were found using flux monitoring, the characteristics of thermogenic methane were identified using carbon isotope tracing, and the relationship between emission by soils and effusion from gas hydrates was determined through correlation analyses of soil-adsorbed hydrocarbons. Seasonal variation of methane emissions are also discussed by considering the emission of bacterial methane, thermogenic methane, and the absorption of methane from the soil active layer. These comprehensive findings provide valuable information for carbon cycle research of wetlands in permafrost regions.

Methane is the second most abundant greenhouse gas in the atmosphere. Since 2007, global atmospheric methane concentrations have increased rapidly at an annual rate of more than 6.8–10 ppb, and the consequent increase in radiative forcing is one of the reasons for a continued rise in global temperatures. Studies have shown that climate warming will lead to the release of carbon from soil pools, resulting in a positive feedback effect. The emission of methane plays a pivotal role in such positive land-based carbon climate feedback mechanisms. As one of the largest carbon pools in nature, wetlands in permafrost regions release 20–72 Tg of methane every year. This release is modulated by methane sources and varies across the seasons. Therefore, studies on the seasonal variations of methane emission sources are of great significance to understanding wetland methane emission processes, investigating wetland carbon cycling, and evaluating regional greenhouse effects.

The Qinghai-Tibet Plateau contains the largest distribution of frozen alpine soil in the world, with a wetland area spanning approximately 130,000 km². Being one of the key ecological safety barriers in Asia, the rate of increase of the atmospheric background methane concentration in the Qinghai-Tibet Plateau is significantly higher than the global average. Recent research has found that global wetland methane emissions increase continuously during the process of climate warming. There are two main theories that explain this phenomenon: (1) elevated temperatures promote soil microbial activity at deeper horizons, thus increasing the environment's carrying capacity for methanogenic microbial communities and increasing methane emissions from microbes; (2) plant activity increases with temperature, thereby influencing the methanogenic microbial community structures and promoting bacterial methane emissions. Compared with conventional peat wetlands, the wetlands in the permafrost regions of the Qinghai-Tibet Plateau possess high-altitude alpine climatic characteristics, which result in higher sensitivities of the wetland ecosystems to temperature changes. In addition, substantial amounts...
of soil organic carbon and thermogenic hydrocarbons in gas hydrates are stored in the Qinghai-Tibet Plateau. These hydrocarbons contribute to methane emission mechanisms that are unique to the region.

Studies on carbon cycling in the wetlands of the Qinghai-Tibet Plateau indicate that elevated temperatures can increase plant and microbial activity, which can lead to increased soil organic carbon content and increased wetland methane emissions. In particular, methane emission bursts have been observed in wetlands during the growing season. Mackelprang et al. attributed the methane emission bursts in wetlands of permafrost regions to methane releases from deep permafrost layers. Lu et al. asserted that the source of methane emissions from wetland surfaces in the Qinghai-Tibet Plateau could be hydrocarbon gases released by subterranean gas hydrates. Sun et al. pointed out that hydrocarbon gases in gas hydrates can effuse towards the land surface through microseepage. According to the hydrocarbon microleakage theory, the hydrocarbon gases in the effusion of gas hydrates may have an impact on the surface carbon circulation system, forming a key source of wetland methane emissions. However, there was lack of evidence for the contribution of the hydrocarbon in gas hydrates to soil methane emissions. Therefore, in studies on wetland methane emissions in the permafrost regions of the Qinghai-Tibet Plateau, the effusion of methane from gas hydrates—which has not been previously explored in detail—must also be considered.

The gas hydrates of the permafrost regions of the Qinghai-Tibet Plateau were first observed in 2008. They are mainly distributed in the pores and fissures of fine-grained sandstones, siltstones, mudstones, and oil shales of the Middle Jurassic Jiangcang Formation, at depths of 133–396 m. The hydrocarbons in gas hydrates are mainly thermogenic, with a dryness coefficient (C1/C2) ranging from 1.3 to 26.0 and a δ13C methane value ranging from −52.7‰ to −35.8‰. As the δ13C content of methane between thermogenic and bacterial gases differs, researchers proposed diagrams to distinguish between methane gas sources. Few studies on methods to distinguish between the sources of wetland methane gas in the Qinghai-Tibet Plateau have been reported.

In order to get an understanding of the source and seasonal variation patterns of methane emissions, in this study, a systematic soil methane cycle geochemical monitoring has been carried out in the Muli area located in the northeastern region of the Qinghai-Tibet Plateau (Fig. 1a). As a typical wetland, the study area is situated 4100–4300 m above sea level. The soil active layer thickness and permafrost thickness are 2.4 and 80 m, respectively. The wetland ecosystem is mainly influenced by freezing and thawing of the permafrost which is governed by seasonal temperature variations. Microbes and thermogenic methane were characterized according to the δ13C content of methane and the seasonal variation patterns of methane emission were discussed based on the geochemical analysis. At present, the methane emissions of the wetland soil in the Qinghai-Tibet plateau have not been investigated in full, and our study provides a novel way of source apportionment for soil methane emissions in permafrost regions.

Results

Methane content of low-level air and soil-adsorbed gas. Table 1 shows the methane content of the low-level air and soil-adsorbed gas samples that were collected. The results (Fig. 2) indicate that the near-surface low-level air data had the smallest dispersion, with a coefficient of variation ranging from 9.62% to 29.00%. The methane concentration of low-level air was significantly higher than that of the regional base air (Fig. 2a). Data on the methane content of near-surface soil-adsorbed gas had the largest dispersion, with a coefficient of variation ranging from 194.37% to 294.43% (Fig. 2b). Data on the heavy hydrocarbon content of near-surface soil-adsorbed gas had a smaller dispersion, with a coefficient of variation ranging from 52.99% to 111.31% (Fig. 2c). The methane content of low-level air and soil-adsorbed gas, as well as the heavy hydrocarbon content (C1-C8) of soil-adsorbed gas exhibited consistent seasonal variation trends, with the highest values in summer, followed by spring and autumn.

The correlation between methane content and heavy hydrocarbon content in surface soil-adsorbed gas can reflect the effusion of thermogenic gases that formed underground. Geochemical analyses of cores from gas hydrate wells in the study area revealed that the gaseous portion of gas hydrates is dominated by methane and also contains a substantial amount of heavy hydrocarbons. Such gases are generated by the large-scale migration of hydrocarbon gases in deep source rocks. During the upward effusion of gases from gas hydrates, methane and heavy hydrocarbons simultaneously effuse towards the surface, thereby directly influencing the hydrocarbon content of near-surface soil-adsorbed gas.

Table 2 shows the results of correlation analyses performed on the collected data. The correlation coefficients between methane content and heavy hydrocarbon content in spring, summer, and autumn were 0.48, 0.17, and 0.88, respectively. Previous studies have reported that bacterial gases have a high methane content and an extremely low heavy hydrocarbon content. The methane content and heavy hydrocarbon content during summer were significantly correlated (R = 0.88), which indicates that effusion from gas hydrates was the main contributor to methane at wetland surfaces. In contrast, the effusion of underground hydrocarbons did not produce a significant influence on the land surface during spring and summer.

Carbon isotope content of methane. Table 3 shows the carbon isotope content of methane for the collected samples. The results (Fig. 3) indicate that samples collected in the spring had the lowest mean carbon isotope value of −57.27‰ Peedee Belemnite (PDB) formation from South Carolina, USA, as well as the largest dispersion with a coefficient of variation of −23.19%. Samples collected in the summer showed a lower methane carbon isotope content (mean value of −52.52‰ PDB) and a smaller dispersion (coefficient of variation of −14.93%). Samples collected in the autumn showed the lowest carbon isotope methane content (mean value of −52.52‰ PDB) and a smaller dispersion (coefficient of variation of −16.25%).

The δ13C1 content of methane and the dryness coefficient of gas hydrates can indicate the source of hydrocarbon gases: δ13C1 < −55‰ PDB indicates bacterially-derived methane, and δ13C1 > −50‰ PDB indicates...
thermogenically-derived methane. The Bernard diagram was used to distinguish between gas sources, the results of which are shown in Fig. 4a. The methane gas sources in the study area exhibited strong polarization during spring. Among the samples collected in spring, 18 were thermogenically-derived methane (43% of all samples) with $\delta^{13}C_1$ contents of $-49.86\%$ to $-38.76\%$ and dryness coefficients of $2.30$ to $18.68$; 12 samples were bacterially-derived methane (29% of all samples), with $\delta^{13}C_1$ contents of $-81.39\%$ to $-60.12\%$ and dryness coefficients of $119.09$ to $2477.27$. The methane sources in summer were similar to those in spring: 18 samples were thermogenically-derived methane (43% of all samples) with $\delta^{13}C_1$ contents of $-48.85\%$ to $-41.16\%$ and dryness coefficients of $3.23$ to $11.64$; 9 samples were bacterially-derived methane (21% of all samples) with $\delta^{13}C_1$ contents of $-64.45\%$ to $-52.99\%$ and dryness coefficients of $218.49$ to $1556.48$. In autumn, the surface methane exhibited a strong thermogenic nature, with 33 samples being thermogenically-derived methane (70% of all samples) with $\delta^{13}C_1$ contents of $-50.00\%$ to $-29.16\%$ and dryness coefficients of $3.24$ to $15.46$; and only one sample was bacterially-derived methane (Fig. 4b).

Compared with the actual gas hydrate samples, the thermogenic samples in the three groups had identical $\delta^{13}C_1$ values and slightly higher dryness coefficients. Results of a gas hydrate simulation experiment indicated that methane exhibits the greatest migration extent during the upward effusion of underground hydrocarbon gases. In contrast, heavy hydrocarbons (C$_2$–C$_5$) tend to enter the hydrate lattices. This fractional distillation process during hydrocarbon effusion leads to an increase in the dryness coefficient of the soil active layer-adsorbed gas. The samples from all three seasons contained outliers characterized by lighter carbon isotopes ($>-55\%$ PDB) and higher dryness coefficients ($>100$) (Fig. 5b), which are generally indicative of bacterially-derived hydrocarbons.

**Seasonal variation of methane sources.** The observed variations in the carbon content of methane reflect the seasonal variations in methane gas sources. As shown in Fig. 5a, the relationship between the carbon isotope contents of spring and summer samples can be generally divided into two parts with correlation coefficient values of $-0.63$ and $0.54$. The P$_1$ zone consists of 16 samples: the spring samples containing bacterially-derived methane with $\delta^{13}C_1$ contents of $-84.11\%$ to $-62.15\%$ PDB, and the summer samples having heavier isotopic...
contents, higher proportions of thermogenically-derived methane, and $^{813}C_1$ contents of –64.45‰ to –41.16‰ PDB. In this zone, the dominant sources of wetland methane gas changed from bacterial sources to bacterial and mixed sources. The P2 zone consists of 16 samples: the spring samples containing thermogenically-derived methane with $^{813}C_1$ contents of –47.15‰ to –38.76‰ PDB, and the summer samples having lighter isotopic contents,
increased proportions of bacterial methane, and δ^{13}C_{1} contents of −64.39‰ to −42.90‰ PDB. In this zone, the dominant sources of wetland methane gas changed from thermogenic sources to bacterial sources. The characteristics of source variations in both zones indicate that changes occurred in both bacterial and thermogenic gas sources during the transition from spring to summer.

Figure 5b shows the relationship between the carbon isotope contents of summer and autumn samples can also be generally divided into two parts with correlation coefficient values of 0.97. The P_{3} zone consists of 16 samples, with the summer samples containing bacterially-derived methane with δ^{13}C_{1} contents of −64.45‰ to −58.17‰ PDB. The P_{4} zone consists of 19 samples, with summer δ^{13}C_{1} contents of −48.85‰ to −41.16‰ PDB and autumn δ^{13}C_{1} contents of −48.85‰ to −41.16‰ PDB, which indicates heavier carbon isotope content. The consistent characteristics of source variations in both zones indicates a decrease in bacterial activity and an increase in thermogenic production during the transition from summer to autumn, resulting in wetland methane gas sources being dominated by thermogenically-derived sources during autumn.

**Methane flux from the upper active layer.** Recent studies have suggested that the upper active layer of soil (0–30 cm) in wetlands is an important methane sink. This absorption can greatly affect the soil methane emission. Table 4 shows the soil-atmosphere methane flux data of the upper active layer obtained from the experiments described in Section 2. Figure 6 shows the annual variation of methane absorption in the soil active layer. The mean monthly methane absorption by soil was −11.47 mg/m^{2}·h^{-1} in March and gradually increased until
April; in June, the methane absorption flux increased rapidly, reaching a maximum value of –52.61 mg/m–2·h–1. In August, the methane absorption flux decreased gradually, with a mean value of –46.26 mg/m–2·h–1. Changes in methane absorption flux were gradual throughout November to December, with mean absorption flux values of –33.01 mg/m–2·h–1 and –36.57 mg/m–2·h–1, respectively. The methane absorption effect of soil was relatively weak in spring, strengthened during early summer, gradually weakened towards the end of summer, and became weak again in autumn and winter. From March to April, both the atmospheric temperature and methane absorption flux increased gradually; from April to June, the atmospheric temperature increased further, prompting the onset of the growing season for wetland organisms.
and causing a rapid increase in methane absorption flux, consistent with results obtained by Turestsky and Vitt\(^{42}\). From June to August, as the atmospheric temperature increased above 0 °C, the permafrost thaw depth peaked and the methane absorption flux gradually decreased. Studies have shown that permafrost thaw leads to changes in soil temperature and wetland water levels\(^{43,44}\), which promotes methane emissions by microbes under anaerobic conditions\(^{45,46}\). Elevated temperatures also increase the activity of methanotrophic microbes in plants and soil at wetland surfaces, resulting in an increase in methane absorption flux and maintenance of the carbon sink function of the upper active layer\(^{18}\). From June to August, the methane absorption flux in the upper active layer exhibited a decreasing trend, which indicates that the increase in wetland methane emissions exceeded the increase in methane absorption by the soil, leading to changes in the carbon source–carbon sink dynamic geochemical balance in wetland soil.

Previous studies suggested that the gas hydrate mining activities did not affect surface greenhouse gas composition\(^{47}\). However, in August, four high methane emission fluxes were observed, with values of 10.40–76.00 mg/m\(^2\)·h\(^{-1}\). The carbon isotope (\(\delta^{13}C\)) content of methane in the free gas at the methane emission sites was between −47.96‰ and −42.22‰ PDB, which corresponds to thermogenic methane. Therefore, the decrease in the methane absorption effect of wetland soil may be caused by increased thermogenic methane emissions. From August to November, the atmospheric temperature decreased below 0 °C and the methane absorption flux gradually decreased to a minimum with this decrease in temperature, returning to the methane absorption level observed for the spring season prior to the start of the next seasonal cycle.

**Discussion**

According to the above results, the seasonal variations of methane emissions can be summarized as shown in Fig. 7. During spring, microbes are the dominant sources of methane. As an increase in temperature promotes biological activity, the methane absorption effects of methanotrophic microbes and adsorption effects of the soil are enhanced, which results in a slow increase in the methane absorption flux of soil during spring (March to April). The combined effects of methane produced by methanogenic microbes and thermogenic methane lead to a lower methane content in low-level air (Fig. 4a).

During the transition from spring to summer, an exponential increase in the scale of microbial communities occurs along with the increase in temperature\(^{48}\). As the structure of methanogenic microbial communities is also influenced by increased plant productivity\(^{19}\), the methanogenic productivity of these microbes increases to a greater extent compared with the increase in the soil methane absorption ability. As the atmospheric temperature in the study area remains below 0 °C in early summer, thawing action in the permafrost is relatively weak, which leads to smaller methane contributions from thermogenic sources and a slow increase in the atmospheric methane concentration.

The average daily temperature, low-level atmospheric methane concentration, and wetland methane emissions reached a maximum during the summer months of June to August. However, previous research has shown that with an increase in temperature, to a certain extent, the upper limit of the environmental carrying capacity of methanogenic microbial communities is reached, which results in an extremely slow increase in methane concentration\(^{48}\). Therefore, the increase in atmospheric temperature does not lead to a rapid increase in wetland methane emissions. In the present study, thermogenic methane emission sites were identified in wetland soil during August through methane flux monitoring and carbon isotope analysis of the soil free gas. From this finding, we deduce that the methane emission bursts from wetlands near the end of the growing season may be attributed to three effects: (1) the adsorbed methane content in wetland soils approaches the saturation point, leading to weaker absorption effects in the soil; (2) as the temperature increases, slow increases occur in the emission of

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**Figure 7.** Seasonal variations of methane gas sources.
bacterially-derived methane; and (3) the depth of thawed permafrost increases, thereby enhancing the effusion of methane from subterranean gas hydrates.

From late summer to autumn and winter, decreases occur in microbial activity, the scale of microbial communities, and the absorption effects of the active layer towards methane as the temperature decreases. This leads to a substantial reduction in the amount of methane produced by methanogens. In addition, as the average daily temperature falls below 0 °C, freezing occurs in the permafrost, which leads to the decreased effusion of methane from subterranean gas hydrates. During autumn and winter, the heavy hydrocarbon content in the surface soil-adsorbed gas increases and methane from subterranean gas hydrates may seep through pathways such as drainage systems or faults, or migrate to the surface through slow diffusion effects, becoming the dominant source of wetland methane within this period.

Conclusions
This study focused on systematic soil methane cycle geochemical monitoring in a typical gas hydrate region in the Qinghai-Tibet Plateau. The measurements presented in this work provide evidence that the effusion of natural gas hydrate underground is a methane source of wetland soil in permafrost regions. The significantly correlation between methane and heavy hydrocarbon content shows the same source of abiotic methane. The heavy value of carbon isotope content of methane in near surface gas indicates that the thermogenic hydrocarbon in gas hydrate has been discharged to the surface. The $\delta^{13}$C-methane ($-47.96\%$ to $-42.22\%$ PDB) in free gas at the methane emission sites also confirmed the existence of thermogenic hydrocarbon effusion. Hence, as shown in previous studies, the absorption of methane in soil active layers is an important factor affecting soil methane emissions.

Nonetheless, the process of methane emission from natural gas hydrates in the Qinghai-Tibet plateau is still not fully understood. With this in mind, the results of this research provide evidence of the effusion of thermogenically-derived methane from natural gas hydrates. These methane emissions may continue to increase as the climate warms and with further development of gas hydrate resources. Research on gas hydrate methane emissions will facilitate the understanding of carbon cycling and may allow for exploration of the mechanisms of soil methane emissions in the permafrost regions of the Qinghai-Tibet Plateau.

Methods
Sample collection and analysis. As a first step in this study, a data collection zone with an area of 3 km$^2$ was established. By applying a grid to the data collection zone, a total of 42 ($7 \times 6$) monitoring sites were set up with a grid interval of 250 m (Fig. 8a). Low-level air samples from the land surface and near-surface soil samples were collected in May, August, and October 2016 and analyzed for methane content, gas chromatography (GC) of
soil-adsorbed hydrocarbons, and carbon isotope content of methane. Soil-methane fluxes from the upper active layer (20–30 cm) at each of the monitoring sites were recorded in March 4–15, April 12–22, June 19–25, August 15–25, November 13–17 and December 3–7 2016.

**Methane content in low-level air and near-surface soil physically adsorbed gas.** Near-surface low-level air samples were collected using the water displacement method. A 400-ml bottle filled with water was placed on the ground surface and air was collected in it via water displacement. Subsequently, the bottle was left to stand for at least 30 min to facilitate adequate exchange between the air within the bottle and low-level air at the ground surface before the bottle was sealed onsite. Secured by the soil van der Waals force, the adsorption-desorption processes were less affected by temperature. The gas samples were collected from the clay or silt layers of the Quaternary coverage area (Fig. 2b) at a depth of 20–30 cm. After removing gravel and grass roots, each sample was placed in a bottle pre-filled with 200 ml of saturated brine. When the liquid level of the saturated brine reached the 400-ml graduation mark, the bottle was sealed, and placed in an inverted position. More details about the physically adsorbed gas sampling were described by Sun et al.17.

The methane content was measured using gas chromatography (GC) (Agilent 7890A, USA). Before sample analysis, 20 mL ± 0.1 uL of standard gas was accurately drawn with a microsyringe and used for equipment calibration to ensure a relative error of less than 3%. After calibration, the six-port valve was heated to 100 °C ± 5 °C, the pre-column was frozen with a liquid nitrogen cold trap, and a carrier gas flow rate of 25 mL/min was used for enrichment via the six-port valve. Then, 50 mL of the gas sample were drawn using a syringe and injected into the injection port. After air separation, the six-port valve was switched to the analysis position and the GC program was initialized for quantitative calculations and plotting of the GC spectra.

**Carbon isotope content of methane in near-surface soil free gas.** The stable carbon isotope content of methane in near-surface soil free gas samples was measured using a G2132-i Isotope Analyzer (Picarro, USA) (Fig. 2c), which utilizes cavity ring down spectroscopy (CRDS) and has a measurement accuracy of less than 0.8‰. The analyzer was equipped with a cylindrical stainless-steel chamber with a cross-sectional area of 0.19 m², an internal volume of 19.2 L, and 1/4-in (6.35 mm) adaptor sleeves for the chamber inlet and outlet. At each sampling site, a hole with a depth of 20–30 cm was excavated, and then, the chamber was embedded immediately and sealed with soil. The chamber was connected to the instrument, and the system was allowed to warm up for 60 min after a cold boot. After adjustments were made to the instrument, readings were collected under a high-accuracy mode at intervals of 5 min for a total duration of 60 min. The measurements were then saved and recorded on a storage card.

**Monitoring of near-surface soil-atmosphere methane flux.** Near-surface soil-atmosphere methane fluxes were measured onsite (Fig. 2d) using a portable soil fluxmeter (WEST Systems WS-L1840, Italy) connected to a cylindrical stainless-steel chamber with a cross-sectional area of 0.06 m² and an internal volume of 4.7 L and a 1/4-in (6.35 mm) adaptor sleeve. A module utilizing tunable diode laser absorption spectroscopy (TDLAS) was installed in the fluxmeter for continuous dynamic monitoring of methane content (ppm). The absolute atmospheric temperature ($T_a$), pressure ($P$), and specific gas constant ($R$) were also recorded during the monitoring period. Using the rate of change in methane content ($F$, ppm/s), the net air inflow ($A$) and the volume ($V$) of the chamber, the surface-atmosphere methane flux ($M$, mol·m⁻²·h⁻¹) was calculated using the following equation (measurement accuracy: 3% of reading, repeatability: 1.5%, methane flux measurement range: 0.5–150.0 mol·m⁻²·d⁻¹):

$$M = F \cdot \frac{86400 \cdot P}{10^5 \cdot R \cdot T_k} \cdot \frac{V}{A}$$

In previous studies, the carbon flux was measured between 8:00 to 10:0019 and 13:00 to 17:00 local standard time50. Since it usually rains/snows in the study area, the methane flux was monitored between 8:00 and 10:00 local standard time. At each monitoring site the air inlet was located in a pre-excavated hole with a depth of 20–40 cm. After sealing the chamber in the hole, the chamber and the instrument were connected and the system was allowed to warm up for 20 min after a cold boot. When the correlation coefficient ($R$) was greater than 0.8, the instrument readings were collected, saved, and recorded on a storage card.

**Data processing and analysis.** All experimental data were considered for analysis; these data were collated using Microsoft Excel 2013 and differences among the seasonal data were compared using their mean and median values. Boxplots exhibiting the distribution characteristics of the data were plotted using Golden Software Grapher V12, with box boundaries being the upper and lower quartiles, and data outside the 1.5 times the inter-quartile range were regarded as outliers. The calculation of correlation coefficients, correlation analysis, and plotting of isotope-dryness coefficient scatter plots were all performed using Microsoft Excel 2013.

**Data availability**
The datasets analyzed during the current study are available from the corresponding author on reasonable request.

Received: 21 August 2019; Accepted: 22 January 2020;
Published online: 05 May 2020
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**Acknowledgements**

The authors would like to express our sincere gratitude to the editors and anonymous reviewers who have put considerable time and effort into their comments on this paper. We would like to thank Editage (www.editage.cn) for English language editing. We are grateful to Prof. Hangxin Cheng and Prof. Chuandong Zhao for their careful reading and useful suggestions, to Mu Kong for his help during field work. We appreciate the valuable suggestions from the anonymous referees. This study is financially supported by the Environmental Monitoring of Terrestrial Natural Gas Hydrate (No. GZHL201400308) and the Geological Survey Program from China Geological Survey (No. DD20190527).

**Author contributions**

S.Z., Writing – original draft; F.Z., Methodology; Z.S., Conceptualization; A.Q., Writing – review & editing; H.W., Project administration; Z.S., Supervision; Z.Y., Investigation; Y.Z., Project administration; S.P., Visualization; P.W., Validation.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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