The origin of methane in the East Siberian Arctic Shelf unraveled with triple isotope analysis

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Abstract. The Arctic Ocean, especially the East Siberian Arctic Shelf (ESAS), has been proposed as a significant source of methane that might play an increasingly important role in the future. However, the underlying processes of formation, removal and transport associated with such emissions are to date strongly debated.

CH4 concentration and triple isotope composition were analyzed on gas extracted from sediment and water sampled at numerous locations on the shallow ESAS from 2007 to 2013. We find high concentrations (up to 500 µM) of CH4 in the pore water of the partially thawed subsea permafrost of this region. For all sediment cores, both hydrogen and carbon isotope data reveal the predominant occurrence of CH4 that is not of thermogenic origin as it has long been thought, but resultant from microbial CH4 formation. At some locations, meltwater from buried meteoric ice and/or old organic matter preserved in the subsea permafrost were used as substrates. Radiocarbon data demonstrate that the CH4 present in the ESAS sediment is of Pleistocene age or older, but a small contribution of highly 14C-enriched CH4, from unknown origin, prohibits precise age determination for one sediment core and in the water column. Our sediment data suggest that at locations where bubble plumes have been observed, CH4 can escape anaerobic oxidation in the surface sediment.

1 Introduction

The Arctic subsea permafrost harbors a very large active carbon pool of similar size to the terrestrial Siberian permafrost reservoir (Shakhova et al., 2010a). Between 12 and 5 kyr before present (BP), the Holocene transgression (Bauch et al., 2001) submerged extensive parts of the Pleistocene age...
terrestrial permafrost in northern Siberia, forming the very shallow East Siberian Arctic Shelf (ESAS; Romanovskii et al., 2005). As a result, the formerly terrestrial permafrost has been continuously exposed to increasing seawater temperature, salt and anoxic conditions (Dmitrenko et al., 2011; Nicolsky et al., 2012), allowing the remobilization of carbon from the Pleistocene reservoirs. The four suggested mechanisms controlling the release of Pleistocene carbon to the ESAS are the deepening of the permafrost level, gas hydrate degradation, coastal erosion and riverine discharge (e.g. Shakhova et al., 2005, 2009, 2010a, b, 2015; O’Connor et al., 2010; Winterfeld et al., 2015; James et al., 2016). Holocene-age carbon originating mainly from coastal erosion and riverine discharge (Charkin et al., 2011; Semiletov et al., 2012; Karlsson et al., 2011, 2016) has accumulated on the ESAS and overlays the Pleistocene age sediment (Vonk et al., 2012, 2014; Feng et al., 2013).

Under anaerobic conditions and depending on its type and quality (Schuur et al., 2013), the remobilized carbon can be used to produce CH$_4$. Microbial CH$_4$ is produced by methanogenesis using carbon dioxide (CO$_2$) or acetate as the main substrates according to the following reactions (Whiticar, 1999).

\[(\text{CO}_2 \text{ reduction}) \quad \text{CO}_2 + 4\text{H}_2 \rightarrow \text{CH}_4 + 2\text{H}_2\text{O} \]

\[(\text{Acetate fermentation}) \quad \text{CH}_3\text{CO}_2^- + \text{H}_2\text{O} \rightarrow \text{CH}_4 + \text{HCO}_3^- \]

In the deep Earth layers, CH$_4$ can also be formed through thermal degradation of organic matter (e.g. Schoell, 1988) and migrate towards the surface. This CH$_4$ is considered thermogenic. A large part of the CH$_4$ formed in the seafloor is removed by anaerobic oxidation with seawater sulfate in sediments (e.g. Reeburgh, 2007; Knittel and Boetius, 2009) or in the water column where CH$_4$ can be consumed by aerobic methanotrophic bacteria under specific nutrient and redox conditions (e.g. Kessler et al., 2011; Mau et al., 2013; SteINle et al., 2015). Each type of CH$_4$ formation and/or removal pathway produces CH$_4$ with a characteristic isotopic signature ($\delta^{13}$C and $\delta$D) depending on the isotopic composition of the substrate and the kinetic isotope effect associated with the respective chemical reaction involved. Microorganisms need less energy to metabolize molecules with smaller bond energy, which leads to discrimination against heavy isotopes. Therefore, CH$_4$ produced by methanogenesis has a lighter isotopic signature than its substrates, but when it is consumed, its remaining reservoir will become more enriched in heavy isotopes (e.g. Whiticar, 1999; Conrad, 2005). Diffusive transport can also cause isotopic discrimination, because lighter isotopologues diffuse faster than heavier ones. However, this fractionation is relatively small ($< 5\%e$; Fuex, 1980; $< 20\%e$; Prinzhofer and Pernaton, 1997; and $3\%e$; Chanton, 2005) compared to the isotopic fractionation associated with methanogenesis ($7$–$95\%e$ for $\delta^{13}$C and $260$–$430\%e$ for $\delta$D) and with CH$_4$ oxidation ($2$–$39\%e$ for $\delta^{13}$C and $66$–$350\%e$ for $\delta$D) (Whiticar, 1999; Holler et al., 2009).

Shakhova et al. (2010b) have shown that CH$_4$ concentrations in the ESAS water were anomalously high (up to 500 nM) compared to CH$_4$ values generally observed in ocean waters ($\sim$5 nM, Damm et al., 2008). Vigorous bubbling events (1.5 to 5.7 bubbles per second) were observed at some sites (Shakhova et al., 2013) as well as seepages of thermogenic CH$_4$ (Cramer and Franke, 2005) indicating that part of the water column supersaturation likely results from a seabed source. The destabilization of gas hydrates is frequently discussed as a CH$_4$ source in this region (e.g. Kvenvolden, 1988; Romanovskii et al., 2005; Shakhova et al., 2010a; Ruppel and Kessler, 2017); however, important gaps exist in the assessment of the quantity and the nature of the CH$_4$ stored or formed in the Arctic seabed (e.g. Ruppel, 2014).

To disentangle the origin(s) of this CH$_4$ anomaly, we measured CH$_4$ concentration, stable isotope composition and (on selected samples) radiocarbon content in sediment and water samples from several winter campaigns and summer cruises from 2007 to 2013 on the ESAS and shelf edge. While stable isotope analyses help identify the chemical pathways involved in CH$_4$ removal and formation processes, radiocarbon measurements give information on the age of the CH$_4$ substrate. The combination of the isotope information thus helps in determining the possible origin(s) of this gas. Determining the stable isotope signatures of the main methane sources in the ESAS also remains crucial to better quantify the CH$_4$ emissions in this region using isotopic- and back-trajectory analysis of atmospheric CH$_4$ (Thornton et al., 2016).

2 Method

2.1 Drilling and sediment sampling

Summer surface sediment drilling and water sampling campaigns were carried out on research vessels, while the winter field campaigns were accomplished using an equipment caravan, which traveled over the sea ice to the drilling locations. In the latter case, casings were drilled through the fast ice into the seabed, allowing dry drilling using a rotary drill with 4 m casing with a newly built URB-4T drilling rig (made in 2011 by the Vorovskii Factory for Drilling Equipment, Ekaterinburg, Russia). Thawed and frozen sediments for each core were subsampled straight after (i.e. maximum a few minutes after) the drilling using ice screws for frozen samples and a heavy plastic syringe-like sampler for thawed samples at 20 cm vertical resolution.

2.2 Gas extraction and measurement in sediments

Sediment subsamples were subsequently immersed in glass vials filled with a saturated sodium chloride solution to drive gases out of solution, and capped with a septum for equilibration in an ultrasonic water bath at a temperature of 20 °C. The gas chromatograph (GC) used to measure CH$_4$ concen-
trations was equipped with two 10-Port gas sampling valves, a 2 m MolSieve 13X column, a 30 m capillary column and a 6 channel PeakSimple data system. A flame ionization detector (FID) was used for concentrations of CH$_4$ < 200 ppm and a thermal conductivity detector (TCD) for concentrations of CH$_4$ > 200 ppm. The GC oven was operated isothermally at 40°C and the maximum detector temperature was held at ≈ 250°C. The carrier gas used was helium. Daily calibration was performed with certified 1.96 and 99.999 ppm CH$_4$ gas standards from Air Liquide, USA. The standard deviation of duplicate analyses (three to five replicates) was < 2%. Reproducibility was ∼ 1% based on multiple standard injections during daily calibrations. The concentration of dissolved CH$_4$ in the water and sediment samples was calculated with the Bunsen solubility coefficient for CH$_4$ (Wiesenburg and Guinasso, 1979) for the appropriate equilibration temperature, pressure and the volume of headspace and water and/or sediment in each vial.

The stable isotope measurements were performed using a continuous-flow isotope ratio mass spectrometry (CF-IRMS) system as described in Brass and Röckmann (2010) and Sapart et al. (2011). Radiocarbon analyses could be performed only on the largest samples (containing more than 20 μg of CH$_4$). In that case, CH$_4$ was preconcentrated and combusted to CO$_2$. The $^{14}$C content of the CO$_2$ was measured by accelerator mass spectrometry (Szidat et al., 2014) using a specific gas inlet (Ruff et al., 2010).

2.3 Gas extraction and measurement from seawater samples

Water samples were collected directly from the Niskin bottles. Gas from seawater samples was extracted using a modified headspace vacuum-ultrasonic degassing method (Schmitt et al., 1991; Lammers and Suess, 1994). The gas released was accumulated in an evacuated burette to measure its quantity and was then transferred into a smaller flask for storage, and analyzed as described in Sect. 2.2.

3 Results and discussion

We present results of CH$_4$ concentrations, stable isotope composition and (on selected samples) radiocarbon content on four shallow sediment cores (<3 m), four deep sediment cores (ID-11, IID-13, IIID-13, VD-13) (down to a maximum depth of 53 m in the Buor-Khaya Bay) and about fifty water samples from four coastal areas of the ESAS: the Lena Delta, the Buor-Khaya Bay, the Dmitry Laptev Strait and the shelf edge (Fig. 1) (see Table S1 for more detail on the sample locations). Because of the harsh field and weather conditions during this campaign, no sediment drilling was possible at the shelf edge; hence only water data are presented for this site. All water and sediment sampling, except for the ID-11 core, was performed at hotspot sites, i.e., at locations where active gas bubbling from the seafloor and high concentrations of dissolved CH$_4$ were previously observed, as discussed in Shakhova et al. (2010a). The location of core ID-11 is therefore referred to as the “non-ebullition site”. This core, as well as the IIID-core, was thawed all the way down (>50 m) while the IID-13 and VD-13 cores were thawed down to 19 and 12 m, respectively. Note that for the two latter cores, sampling was continued through the deeper frozen sediment to 30 and 35 m respectively. For more details on the lithology, the cryostructure and the sediment properties, see the Supplement, Sect. S1 and Figs. S1–S4.

3.1 CH$_4$ formation pathways in the sediment

Depth profiles of CH$_4$ concentration, stable isotope composition ($\delta^{13}$C and $\delta$D) and the radiocarbon content (in percentage of modern carbon, pmC) are presented in Fig. 2. In both hotspot and non-ebullition cores, CH$_4$ concentrations are far above values observed in the water column, and CH$_4$ is strongly depleted in heavy stable isotopes in all sediment cores. CH$_4$ in the hotspot cores IID-13, IIID-13 and VD-13 is more depleted in D and slightly more enriched in $^{13}$C than in the non-ebullition core. These differences can be caused by the distance of the drill sites from the coast, the amount of time each site has been inundated and the differences in lithology (Supplement, Sect. S1). These factors will play a role in the substrate availability (Karlsson et al., 2011, 2016; Tesi et al., 2014, 2016). We will focus the discussion on the origin of the substrate(s) for each core below.

The expected stable isotope signatures of the three potential CH$_4$ formation pathways in marine sediment (e.g. Whiticar, 1999), CO$_2$ reduction, acetate fermentation and thermal degradation of organic matter, are depicted together...
with our water and sediment stable isotope data in a dual isotope plot (Fig. 3). Overall, the deep sediment core data (diamonds) fall in between the isotope source signatures of the two main microbial CH₄ formation pathways: carbonate reduction and acetate fermentation. These atypical stable isotope signatures could imply that CH₄ is formed by a mixture of both microbial pathworks and/or by using different substrates from the ones considered in Whiticar (1999). It is unlikely to be explained by physical alteration (e.g. diffusion, gravitational settling) because these processes would result in equal fractionation for the CH₃D and CH₃C isotopologues.

For the non-ebullition core ID-11, most of the δ¹³C values are typical (though on the light end side) of the reduction of carbonates, but about 2/3 of the samples show δD values that are considered too low (down to about −60 ‰) for such a pathway. The most enriched δD data correspond to the top of this core and are discussed in Sect. 3.2. For this core, salinity measurements (from 20 PSU at the surface to 13 PSU at depth) indicate the presence of interstitial seawater all the way down the core. When the seawater sulfate enters the marine sediment, it provides sulfate-reducing bacteria with the electron acceptor they need to out-compete methanogens for acetate (Lessner, 2009). This indicates that for this core, in situ (i.e. at the depth where the samples were taken) acetoclastic CH₄ formation may be suppressed, despite an abundance of organic material. CO₂ and water therefore remain the most likely non-competitive substrate for methanogens if CH₄ formation would occur in the thawed permafrost. In that case, the very low δD values can be due to (1) a mixture of carbonate-reduced (formed in situ or not) and acetoclastic (migrating vertically or horizontally) CH₄ or (2) the use of isotopically depleted hydrogen substrate for CH₄ formation by carbonate reduction. On the dual isotope plot (Fig. 2), the area of the carbonate reduction pathway considers modern seawater as water substrate for carbonate reduction. However, the meltwater present in subsea permafrost originates from buried meteoric ice with much more depleted δD(H₂O) signatures. Chanton et al. (2006) and Brosius et al. (2012) reported values for δD(H₂O) of −135 ± 25 ‰ and −220 ± 30 ‰, respectively in old Arctic permafrost. This is about 200 to 105 ‰ more depleted in deuterium than modern Arctic seawater (Friedman et al., 1964). We suggest that methanogens present in the thawing permafrost (Koch et al., 2009) use and/or have used such depleted permafrost meltwater or unfrozen porewater as a hydrogen source to form CH₄ with low δD values as it is observed in the non-ebullition core.

For the hotspot cores IID-13 IIID-13 and VD-13, the δD values are characteristic of acetate fermentation, but the δ¹³C signatures are about 30 ‰ more depleted in ¹³C in comparison to what has been measured previously from this pathway (e.g. Whiticar, 1999; Walter et al., 2008). This depletion in ¹³C must originate from (1) the addition of carbonate-reduced CH₄ to an acetoclastic pool and/or (2) the recy-
Our cores were relatively constant and not correlated with the concentration, but this was not observed: the CH$_4$ concentrations in our cores were relatively constant and not correlated with the $\delta^{13}$C values (Fig. 4). For these cores, because of the harsh conditions in the field, no reliable sulfate and salinity profiles could be retrieved, so unfortunately no sulfate data are available to support the interpretation.

The $^{14}$C content of CH$_4$ from the hotspot cores covers a range from 0.79 to 3.4 pmC corresponding to a radiocarbon age of 26 to 39 kyr BP (Fig. 2). This indicates a carbon substrate of Pleistocene age. For the ID-11 non-ebullition core, $^{14}$C values are unexpectedly high and vary from 87 pmC (radiocarbon age = 1 kyr BP) to 2367 pmC (Fig. 2), which represents a substantial enrichment above the natural background. The same applies to water samples from the shelf edge. Note that levels close to 100 pmC indicate modern values. Even samples that had been affected by the nuclear bomb testing in the 1950s and 1960s would show levels below 200 pmC; thus $^{14}$C values > 200 pmC cannot be caused by known natural processes. As discussed in the Supplement Sect. S2, local anthropogenic nuclear contribution, e.g. from nuclear waste buried in the coastal permafrost, is the most likely explanation for these elevated radiocarbon levels. The drilling location is shallow (12.5 m) and very difficult to reach and hence waste burial is very unlikely to have occurred directly in this area. Moreover, the highest contamination is observed at 30 m depth in the sediment, showing that it may not originate from the surface. Our first suggestion is that this anthropogenic contamination has been laterally transported in the pore water of the thawing subsea permafrost in the form of CH$_4$ or of one of its precursors (e.g. dissolved inorganic carbon) from the coastal terrestrial permafrost to our drilling site (see Supplement Sect. S2 for more detailed). More data (e.g. of other radionuclides) would be essential to confirm this assumption.

The shallow sediment samples from hotspot sites have $^{14}$CH$_4$ values from 3 to 88 pmC (radiocarbon age = 1–26 kyr BP), showing the presence of old CH$_4$ in surface sediment of relatively modern age and thus confirming the migration of old gas from deeper layers towards the surface. Note that the overall low content of organic carbon (< 2.3 %) with a high fraction of lignin (Bröder et al., 2016; Vonk et al., 2014) in the surface sediment (Fig. 5), and the likely presence of sulfate, would severely inhibit CH$_4$ formation in the marine layer; hence in situ methanogenesis there is highly unlikely.

We conclude that the CH$_4$ present in the surface thawed subsea permafrost is formed mainly microbially. For the non-

**Figure 4.** CH$_4$ concentration versus stable isotope plots. Water sampling (triangles), sediment cores (diamonds). Summer sampling (close symbols) and winter sampling (open symbols). Buor-Khaya Bay (purple, ID-11: non-ebullition site and IID-13, IIDD-13 and VD13 hotspot sites), Dmitry Laptev Strait (red and orange), Lena Delta (light blue) and Shelf Edge (yellow) (see Fig. 1 for detailed locations and Table S1 for bathymetric information).

**Figure 5.** Close-up of the CH$_4$ concentration, stable isotope and other biogeochemical data of the surface of the non-ebullition sediment core ID-11, from the Buor-Khaya Bay. Red shaded area corresponds to the marine sediment deposited during the Holocene transgression and the grey shaded area corresponds to the thawed permafrost layer. The black dotted line corresponds to the depth where CH$_4$ oxidation starts to occur.
ebullition core, our observations imply that CH₄ is at least partly not formed in situ in thawed subsea permafrost, but that it migrates vertically or laterally to the surface of the partially thawed ESAS subsea permafrost. For the hotspot cores, which are closer to the shore and more recently inundated (Table S1), most of the methane present is of acetoclastic origin and formed with Pleistocene carbon remobilized in the thawing subsea permafrost.

3.2 CH₄ removal pathways in the sediment

The ID-11 non-ebullition site was the only coring location where no active bubbling was observed from the surface sediment. Here, the top 5.8 m consists of a thick silty-clay layer (Fig. S1) of marine origin as indicated by the higher salinity and silica concentrations (Fig. 5), typical of a marine environment enriched in diatoms. The increase in sulfate concentration, together with the strong CH₄ concentration decrease and the isotopic enrichment in both Δ¹³C and ΔD towards the sediment surface, indicates that most of the CH₄ diffusing through this thick Holocene marine layer is removed by anaerobic oxidation with sulfate in the surface sediment before reaching the water column.

This marine layer may also act as a physical barrier preventing gas migration towards the surface directly. The increase in CH₄ concentration from 9 to 5.8 m depth without strong isotopic shifts (Fig. 5) and the acoustic data (Fig. 6) show that gas accumulates under this less permeable layer. Part of this gas might migrate laterally and be released to the water at locations where the marine clay layer is thinner or absent. The isotopic signatures of the CH₄ in the pore water of the hotspot cores do not show isotopic fractionation toward the surface (Fig. 2). At these sites, ebullition processes may disturb the sulfate-reducing layer and advection may occur. This would reduce the amount of CH₄ subject to anaerobic oxidation (only dissolved CH₄ is accessible for methanotrophic organisms) and allow direct gas release to the water column.

Overduin et al. (2015) have reported CH₄ concentration and δ¹³C values measured on one sediment core drilled in the Buor-Khaya Bay. The carbon isotopic signature of that core was typical of acetate fermentation in the frozen part of the core, but they observed a strong enrichment in Δ¹³C associated with a decrease in CH₄ concentration directly above the ice-bonded permafrost. They concluded that CH₄ was strongly oxidized in the thawed subsea permafrost before reaching the water column. Our dataset suggests that the Overduin et al. core is not typical for the entire region as we did not observe similar enrichment in either ΔD or Δ¹³C associated with a decrease in CH₄ concentration at the ice-bonded permafrost table for the partly frozen cores IID-13 and VD-13 (Figs. 2, S2 and S4).

3.3 CH₄ in the water

Compared to the sediment samples, CH₄ in the water samples is more enriched in heavy isotopes. The highest CH₄ concentrations in the water column are observed close to the seabed and at the surface in the presence of sea ice (Fig. 2a blue triangles). The Δ¹⁴C values of water samples are between 83 and 9560 ppmC (radiocarbon age = 2 kyr BP) to strongly enriched above natural present-day values) (Fig. 2d) (Supplement Sect. S2). For the water samples we only encountered the highly enriched Δ¹⁴CH₄ values at the shelf edge. As demonstrated by the Δ¹⁴CH₄ data in the non-ebullition core ID-11, this anomaly likely originates from anthropogenic contamination in the sediment. Hence, we suggest that this signature may be diluted over the shelf but becomes indiscernible at locations where a strong release of old CH₄ from the sediment occurs. This could explain the broad range of ppmC values observed in the water column.

Several scenarios may explain the difference in stable isotope signatures between the water samples and sediment samples. The first assumes a mixture of microbial CH₄ with a source that is more enriched in heavy isotopes. This source could be either a water source or thermal degradation of organic matter in the Earth’s deep crust. In the marine environment, CH₄ could in principle be produced at the pycnocline, where natural differences of water density create a “fluid bottom”, on which organic particles and pellets could accumulate as substrate for in situ methanogenesis (Damm et al., 2008; Karl et al., 2008; Sasakawa et al., 2008). In the ESAS, the pycnocline is very shallow, and at the location of sampling, low primary production is expected because of darkness and ice cover in the winter and because of the little available sunlight in the summer due to the high solar zenith angles and the very turbid waters (Semiletov et al.,
They concluded that the CH$_4$ contained in the rich waters of the river was, for the most part, not reaching the marine waters, but that it was released by diffusion into the atmosphere before reaching the bay. A large water source is therefore unlikely to explain the CH$_4$ saturation we observe in the ESAS coastal waters.

Thermogenic emissions from the sediment are possible, especially from the fault zone near the shelf edge where we find strong heavy isotope enrichment in the water. While we have not measured any CH$_4$ with a thermogenic stable isotopic signature in our deep sediment cores from the continental shelf, it could be present in the sediments of the shelf edge (which we were unable to sample due to rough field conditions). Moreover, no measurements could be performed directly on gas bubbles (because of the low probability of trapping bubbles in the Niskin bottles during sampling), which at the shelf edge might partly originate from thermal degradation of organic matter.

The difference between the water and sediment samples may also result from substantial oxidation of the CH$_4$ emitted from the deep sediment. Such a process should involve enrichments in D and $^{13}$C associated with a decrease in CH$_4$ concentration. This pattern is only observed for the winter water samples of the Lena Delta (Fig. 4, blue open triangles) where CH$_4$ trapped under the sea ice could be removed by aerobic oxidation. All other water data were collected in the summer and do not show any clear isotopic enrichment correlated with concentration decrease. This could be explained by the continuous addition of CH$_4$ from the sediment and its direct diffusion from the water into the atmosphere in the summer, especially during storms (Shakhova et al., 2013). These processes as well as water-column mixing could mask any oxidative isotope signature.

In the winter, CH$_4$ likely accumulates under the sea ice where the bubble and dissolved phases could equilibrate and aerobic oxidation could occur, while in the summer the gas bubbles will directly reach the atmosphere. In the sediment, gas bubbles have time to equilibrate with pore water, especially when the gas is trapped under relatively impermeable sediment, e.g. the Holocene marine silty-clay layer. Therefore, we assume that in the sediment, the pore water can be in equilibrium with the gas bubbles, while we suggest that in the summer the seawater bubbles may travel too rapidly to reach an isotopic equilibrium with the dissolved gas and to be oxidized. This means that the CH$_4$ isotopic signature of the gas bubbles may not strongly affect the CH$_4$ dissolved in seawater, which could also explain the difference observed between the water and sediment stable isotopes.

4 Conclusion

Our triple isotope dataset of CH$_4$ from the sediment and water of the shallow ESAS reveals the presence of CH$_4$ of microbial origin formed on old carbon with unexpectedly low stable carbon ($^{13}$C as low as $-108\_\text{‰}$ and hydrogen ($^{2}$D as low as $-350\_\text{‰}$) isotope signatures down to about 50 m under the seabed in the thawed permafrost. These data demonstrate that at locations where a thick marine clay layer is present, this CH$_4$ is partially oxidized before reaching the seawater. However, at locations where ebullition was observed from the seabed, no oxidation was identified in the stable isotope surface sediment profile. In that case, and considering the very shallow water column (< 10 m) in this area, this microbial gas will likely reach the atmosphere when sea ice is absent. Our results show that thawing subsea permafrost of the ESAS emits CH$_4$ with an isotopic signature that cannot be easily distinguished from Arctic wetland emissions when looking only at stable isotope data. This similarity might complicate recent efforts to quantify Arctic CH$_4$ source strengths on the basis of isotopic- and back-trajectory analysis of atmospheric CH$_4$. Further in situ work is necessary – specifically on the isotopic composition of CH$_4$ in gas bubbles that reach the atmosphere – to better quantify the contribution of the ESAS to the global methane budget.

Data availability. The data presented in this paper are available at https://www.projects.science.uu.nl/atmosphereclimate/Data.php.

The Supplement related to this article is available online at doi:10.5194/bg-14-2283-2017-supplement.

Author contributions. Célia J. Sapart, Natalia Shakhova, Thomas Röckmann, Joachim Jansen, Sönke Szidat, Igor Semiletov, Jean-Louis Tison and Matthias Egger worked on the scientific interpretation and wrote the manuscript. Natalia Shakhova and Igor Semiletov planned the research and organized the multiyear fieldwork campaigns. Carina van der Veen, Célia J. Sapart, Sönke Szidat and Joachim Jansen performed the isotopic analyses. Igor Semiletov, Denis Kosmach, Oleg Dudarev, Valentina Sergienko, Anatoly Salyuk and Vladimir Tumskoy performed the water sampling, sediment drilling, headspace preparation and CH$_4$ concentration measurements on the field.

Competing interests. The authors declare that they have no conflict of interest.

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