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Kev Points:

- Summer CH₄ release from seabed sediments west of Svalbard substantially increases concentrations in the ocean, but not in the atmosphere
- The modeled flux is constrained to a maximum of 2.4 to 3.8 nmol m $^{-2}$ s $^{-1}$ compatible with the observed atmospheric CH $_4$ from 20 June to 1 August 2014
- Any ocean-atmosphere flux of the CH₄ accumulated beneath the pycnocline may only occur if physical processes remove this dynamic barrier

Supporting Information:

• Supporting Information S1

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Extensive release of methane from Arctic seabed west of Svalbard during summer 2014 does not influence the atmosphere

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Abstract We find that summer methane (CH₄) release from seabed sediments west of Svalbard substantially increases CH_4 concentrations in the ocean but has limited influence on the atmospheric CH_4 levels. Our conclusion stems from complementary measurements at the seafloor, in the ocean, and in the atmosphere from land-based, ship and aircraft platforms during a summer campaign in 2014. We detected high concentrations of dissolved CH_4 in the ocean above the seafloor with a sharp decrease above the pycnocline. Model approaches taking potential CH_4 emissions from both dissolved and bubble-released CH_4 from a larger region into account reveal a maximum flux compatible with the observed atmospheric CH_4 mixing ratios of 2.4–3.8 nmol m⁻² s⁻¹. This is too low to have an impact on the atmospheric summer CH_4 budget in the year 2014. Long-term ocean observatories may shed light on the complex variations of Arctic CH_4 cycles throughout the year.

1. Introduction

The important greenhouse gas methane (CH₄) has large natural sources vulnerable to climate change [Ciais et al., 2013; Myhre et al., 2013; Portnov et al., 2016]. The causes of the recent global average growth of ~ 6 ppb yr⁻¹ since 2007 in atmospheric CH₄, including a marked Arctic growth event in 2007, remain unclear [Nisbet et al., 2014; Kirschke et al., 2013]. Decomposing methane hydrates (MHs) in marine sediments along continental margins is potentially a large natural source [Ruppel, 2011]. How much of the CH₄ stored or formed by biogenic processes in the Arctic subsea that escapes to the atmosphere remains an open question. Large CH₄ gas escape from the shallow seabed to the ocean column has been reported from East Siberian Arctic shelves (ESAS), particularly during storms [Shakhova et al., 2014] and from the Laptev and Kara Seas [Shakhova et al., 2010; Portnov et al., 2013]. Very high fluxes of CH₄ from subseabed sources to the atmosphere have been reported for the ESAS [Shakhova et al., 2010, 2014], with flux values of \sim 70–450 nmol m⁻² s⁻¹ under windy conditions, with a postulated average total area (extrapolated) source magnitude of $17\,\mathrm{Tg}\,\mathrm{yr}^{-1}$ representing 3% of the global budget to the atmosphere. However, on the contrary it was recently found that the ESAS region only emits from 0.5 to 4.5 Tg yr⁻¹ [Berchet et al., 2016]. Based on continuous atmospheric observations, there are hundreds of gas plumes observed in the water, suggestive of gas release northwest off Svalbard. Along the West Svalbard continental margin, extensive gas bubbling from the seafloor has been observed in shallow water at 90-400 m depth [Knies et al., 2004; Westbrook et al., 2009; Rajan et al., 2012; Sahling et al., 2014; Veloso et al., 2015; Graves et al., 2015; Steinle et al., 2015; Smith et al., 2014; Portnov et al., 2016, this work] outside of today's gas hydrate stability zone [Panieri et al., 2016]. It is unknown how much of the CH₄ flux from the marine sediments in this region ultimately reaches the atmosphere [Fisher et al., 2011], either through bubbles or flux of dissolved CH₄.

The amount of CH₄ stored within gas hydrates, or as dissolved and free gas, north of 60°N is uncertain. Estimates as high as 1200 Gt have been reported [*Biastoch et al.*, 2011]. Some hydrate deposits may be on

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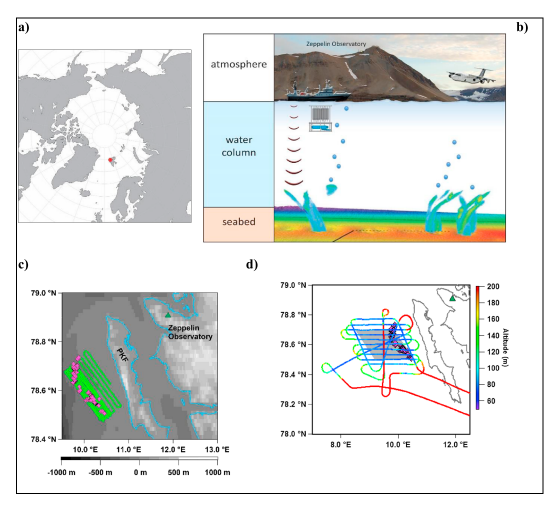


Figure 1. Field campaign and measurement platforms at the seafloor, in the water column, and in the atmosphere west of Svalbard in June-July 2014. (a) The location of the measurement area marked in red west of Svalbard. (b) Illustration of the field activity from 23 June to 2 July 2014 (not to scale). Seeps on the seafloor, represented here by swath bathymetry, release gas bubbles that rise through the water column. The Research Vessel Helmer Hanssen detected gas bubbles and collected water samples at various depths and provided online atmospheric CH₄, CO, and CO₂ mixing ratios and discrete sampling of complementary trace gases and isotopic ratios. The Facility of Airborne Atmospheric Measurements (FAAM) aircraft measured numerous gases in the atmosphere, and an extended measurement program was performed at the Zeppelin Observatory close to Ny-Ålesund. (c) Detailed map of the area of intense shipborne measurements. The ship track (green line) covers an Arctic shelf region, ~80-200 m depth, as indicated by bathymetric data west of Prins Karls Forland (PKF), an area with numerous observed flares [Westbrook et al., 2009; Sahling et al., 2014, and this work, shown as pink symbols]. The location of the Zeppelin Observatory is shown (green triangle), ~50 km from PKF. (d) Flight track over the same region on 2 July; altitude is given by the color scale, and the area used for the flux calculation based on flight data is shown in grey.

the verge of instability due to ocean warming, leading to a debate whether CH₄ release could trigger positive feedback and accelerate climate warming [Archer, 2007; Isaksen et al., 2011; Ferré et al., 2012]. There have been very few studies aimed at detecting and quantifying the potential atmospheric enhancement of this oceanic source around Svalbard and estimating the fluxe contributions. The West Svalbard continental margin is warmed by the northward flowing West Spitsbergen Current, the northernmost limb of the Gulf Stream. There has been an increase in the bottom water temperature in this area of 1.5°C [Ferré et al., 2012] over the last 30 years, while the atmosphere has warmed by as much as 4°C since the early 1970s [Nordli et al., 2014]. Continued warming in this region is expected [Collins et al., 2013]. Consequently, it is crucial to determine whether, and how, CH₄ from the shallow shelf located close to a stable gas hydrate zone on the upper continental margin reaches the atmosphere at present and how this might change in the future. To investigate this, we have conducted an intensive atmospheric and oceanographic survey (Figure 1) in an area with a known high density of hydroacoustically detected gas flares (indications of bubbles in echograms) west of

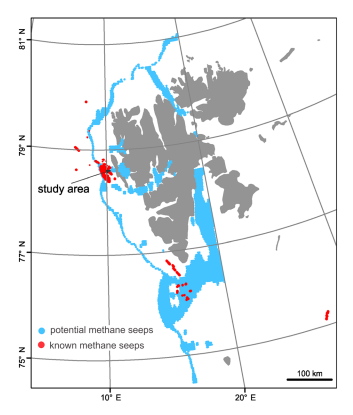


Figure 2. The identified (red) and potential (blue) seep locations around Svalbard as calculated by methane hydrate stability modeling.

Prins Karls Forland, Svalbard, from 23 June to 2 July 2014, with atmospheric measurements continuing to 1 August. We investigated whether there was an atmospheric enhancement and impact during summer time. The measurements were used in combination with three different models to provide independent top-down flux constraints, also taking into account potential emissions from larger areas outside the focused campaign region for the period.

2. Data and Methodology 2.1. Field Platforms, Measurements, and Data

An overview of the area together with the complementary measurement platforms is presented in Figure 1. The research vessel (R/V) Helmer Hanssen was equipped with instruments to analyze water samples from the sea surface down to the seabed and to monitor CH₄ atmospheric mixing ratios from 20

June to onward. A single beam echo sounder constantly recorded flares in echograms; flares represent locations where bubbles are released from the seafloor which rise through the water column [Veloso et al., 2015] and where we expect high-dissolved CH₄ concentrations. Figure 1c shows the ship's route during 24–27 June 2014, together with identified gas flares. Aircraft measurements during the campaign were performed as low as ~15 m above the ocean, covering a larger area than the ship, for a short time (flights were around 4 h in duration). Figure 1d shows the "Facility of Airborne Atmospheric Measurements" (FAAM) aircraft path and height on 2 July 2014 in the area and the location of the flares identified (see Pitt et al. [2016], O'Shea et al. [2013], and Allen et al. [2011] for details of the aircraft and instrumentation). Finally, measurements of the atmospheric composition at the nearby Zeppelin Observatory include continuous CH₄ measurement and daily sampling of CH₄ isotopic ratios (see Table S1); Figure S1 in the supporting information shows the locations. A description of all instruments and methods employed is included in the supporting information. Table S1 gives an overview of the instruments from all platforms involved.

2.2. Model Tools for Data Analysis and Top-Down Flux Estimations

Potential CH₄ seep locations around Svalbard were determined by MH stability modeling. The MH stability model (CSMHYD program) [Sloan and Koh, 2008] was used taking bottom water temperatures World Ocean Database 2013 [Boyer et al., 2013] and sediment thermal gradients (Global Heat Flow Database) from around Svalbard as input parameters. Locations, where the hydrate stability zone outcrops at the seabed, are considered to be potential CH₄ seep locations. These locations were supplemented with all known CH₄ seeps [Sahling et al., 2014; Panieri et al., 2015, this work]. The modeled potential methane seep locations and known methane seeps are illustrated in Figure 2 as light blue and red dots, respectively.

In order to estimate CH₄ fluxes from the modeled seep area (blue in Figure 2) and identified CH₄ seep areas (red in Figure 2), we used three different independent atmospheric models: (1) the Lagrangian particle dispersion model FLEXPART [Stohl et al., 2005], (2) the global chemical transport model Oslo CTM3 [Søvde et al., 2012; Dalsøren et al., 2016], and (3) a Lagrangian mass balance box model [Karion et al., 2013; O'Shea et al., 2014]. See section S3 for details about models and simulations.

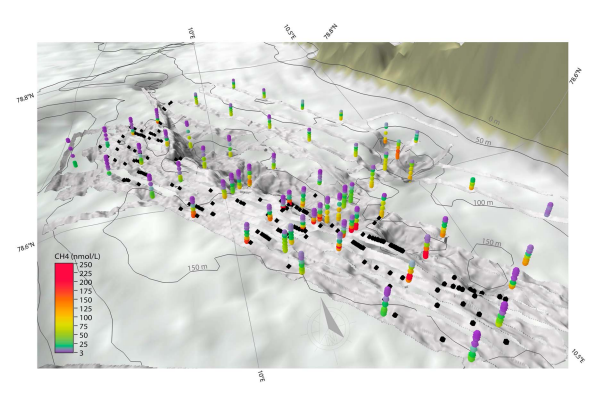


Figure 3. CH₄ concentrations from a hydrocast survey offshore of Prins Karls Forland. The first three bottles were taken 5, 15, and 30 m above the seafloor, and the last three bottles were taken 10, 20, and 30 m below the sea surface. The rest of the samples were spread equally in the water column depending on the bottom depth. CH₄ concentrations in the ocean are illustrated by colored dots (scale on the bottom left in nmol L⁻¹). Black dots indicate the location of the gas flares. Isobaths are from International Bathymetric Chart of the Arctic Ocean version 3 grid and the superimposed higher resolution bathymetry is from the multibeam survey performed during the R/V Helmer Hanssen cruise; data were recorded over the period 25 June to 1 July 2014.

3. Results and Discussion

3.1. Observations in the Ocean and Atmosphere

We present the results following the methane migration path from the seafloor through the water column to the lowermost atmosphere close to the sea surface (ship) and higher up using flight data covering a larger area. Figure 3 illustrates the dissolved CH₄ concentrations sampled over the investigated area. Elevated concentrations were found around the most extended cluster of flares, and the CH₄ distribution shows a rapid change at about ~50 m water depth, with the highest dissolved CH₄ concentrations near the seafloor ~150 m depth. Little CH₄ is found above the pycnocline (boundary where the density gradient is greatest, affected by temperature and salinity), but sea surface CH₄ concentrations are still oversaturated with respect to atmospheric concentrations in a few places eastward, close to the shore

The sea surface CH₄ ocean concentrations (Figure 4a) and the atmospheric mixing ratio measured by both the ship (Figure 4b) and the aircraft (Figure 4c) show very similar patterns. In the surface water CH₄ was generally <8 nmol L⁻¹ (Figure 4a) with a median of 4.8 nmol L⁻¹. A maximum of 26 nmol L⁻¹ was found near the shore, where no gas flares are found in the vicinity (Figures 4a and 4b). The elevated surface water CH₄ concentrations coincide with a small increase (<2 ppb) of atmospheric CH₄ mixing ratio detected by the ship. This slightly elevated CH₄ close to the shore is probably not due to CH₄ released from the seafloor/seeps. Figure 3 and Figure 4 show that the bottom CH₄ concentrations are low in this coastal area. A simultaneous decrease in salinity suggests the intrusion of methane-enriched fresher water [Damm et al., 2005] near the surface increasing the dissolved CH₄ concentrations in this particular area.

A 6 km transect was sampled twice in 1 week by the R/V Helmer Hanssen to monitor rapid variations of oceanographic conditions and their effects on the dissolved CH₄ distribution. The maximum bottom water CH₄ concentration doubled in 1 week from 200 to 400 nmol L⁻¹ (see Figures 4d and S2), while bottom water temperatures remained relatively stable. At the same time, the concentrations above the pycnocline and at sea surface remained relatively stable and low (4–11 nmol L $^{-1}$ and \sim 10 nmol L $^{-1}$ in the surface water on 24 June and 1 July, respectively).

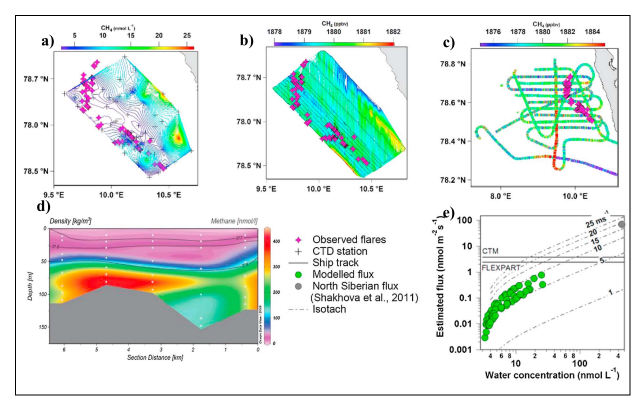


Figure 4. Comparison of CH_4 variations in the ocean and atmosphere west of Svalbard and corresponding CH_4 flux to the atmosphere. (a) Contour plot of near-surface CH_4 concentration (color scale) at ~10 m depth in the water column. CH_4 was measured by oceanographic conductivity-temperature-depth (CTD) stations (crosses) west of Prins Karls Forland (PKF). Observed flares are shown by pink markers. (b) Contour plot of atmospheric CH_4 mixing ratio in parts per billion measured aboard R/V Helmer Hanssen (color scale). Ship track is shown by black line; flares are shown by pink markers. (c) CH_4 measured by the FAAM aircraft; flares are shown by pink markers. (d) CH_4 concentration in the water column along a transect of CTD stations taken on 1 July 2014 showing a clear stratification of water masses with the pycnocline near 50 m water depth. Density is shown as black contours. (The transect location offshore of Prins Karls Forland is shown in Figure S2b). (e) CH_4 flux to the atmosphere at each CTD location as a function of ocean CH_4 concentration according to a diffusive model (green points). Flux previously modeled off Northern Siberia during stormy weather [Shakhova et al., 2014] is given by the grey point. Dashed lines show the model flux at different isotachs (lines of constant wind speed), assuming constant salinity and temperature (averaged over the sampling period used). Horizontal lines show the maximum possible flux constrained by the atmospheric measurements from the ship, according to FLEXPART and OSlo CTM3 models. FLEXPART and CTM constraints are for the atmospheric sampling period 20 June to 1 August and will vary with weather patterns.

This is in agreement with changes reported by *Steinle et al.* [2015] for bottom water and sea surface water. This change in concentration can be explained either by slower advection during the later observations or that the water was previously CH₄ enriched by an emission burst from one or several nearby seep sites. Gas bubble dissolution modeling from a previous study in the deeper area to the west of our study area estimated that 80% of the bubble-released CH₄ is dissolved below the summer pycnocline, and the remaining CH₄ is transported northward where it is most likely oxidized by methanotrophic bacteria [*Gentz et al.*, 2014; *Steinle et al.*, 2015]. A similar conclusion came from a box modeling result of dissolved CH₄ indicating that ~60% of CH₄ released at the seafloor becomes already oxidized before it reaches the overlying surface waters [*Graves et al.*, 2015]. Although our single beam echo sounder studies show bubbles reaching the sea surface, very little CH₄ remains in such bubbles by the time they reach the surface [*Greinert and McGinnis*, 2009].

We compared data from the R/V Helmer Hanssen to those from the Zeppelin Observatory for the period from 20 June to 1 August. The CH₄ mixing ratio measured aboard the ship during the measurements off Prins Karls Forland agrees well with those recorded by the Zeppelin Observatory, as does the isotopic ratio (see supporting information Figure S3). Our measurements above the flares were not influenced by long-range transport of methane-enhanced air masses from lower latitudes, as this would have produced noticeable transient enhancements in CH₄, as exemplified in Figure S3.

3.2. Flux Estimates From Ocean to Atmosphere in the Svalbard Region During Summer

We estimate the median ocean-atmosphere CH₄ flux based on observations in the ocean, in addition to three top-down constrains of the flux employing three independent models and the atmospheric measurements.

Maximum Flux Possible Constrained by the
MethodologyAtmospheric Observations (nmol m $^{-2}$ s $^{-1}$)

3,	•	
FLEXPART ^b top-down backward modeling	2.4 ± 1.4	
Oslo CTM3 ^c top-down forward modeling	3.8 ± 1.4	
Lagrangian mass balancing—FAAM) ^d , top down, exploring upwind/	14.1	
downwind variations		

^aThe potential flux region is shown in Figure 2 and employing atmospheric observation from *Zeppelin* and *Helmer Hanssen* over the period 20 June to 1 August 2014.

We estimate a median ocean-atmosphere CH_4 flux of $0.04\,\mathrm{nmol\,m^{-2}\,s^{-1}}$ (σ = 0.13) from data at each conductivity-temperature-depth (*CTD*) station using an ocean-atmosphere gas exchange function [*Wanninkhof et al.*, 2009] (Figure 4e). The maximum flux at the CTD stations is $0.8\,\mathrm{nmol\,m^{-2}\,s^{-1}}$ which occurred when both dissolved CH_4 concentrations and wind speeds were high, 25 nmol L^{-1} and 9 m s⁻¹ respectively. This model only considers air-sea exchange via diffusion of dissolved CH_4 and not the contribution of bubbles of gas reaching the surface. Figure 4e) shows the estimated flux at different wind speeds, assuming constant salinity and temperature (average from the campaign). Wind speed has a large effect: an increase from 5 to $10\,\mathrm{m\,s^{-1}}$ increases the modeled flux by almost an order of magnitude. The atmospheric CH_4 air mixing ratios aboard the R/V *Helmer Hanssen* and at *Zeppelin* before, during, and after the ship-based measurements off Prins Karls Forland were very similar, with small variations (Figure S3). Hence, the CH_4 air mixing ratios above active seep areas were representative of wider regional atmospheric concentrations, with no elevated levels or transient large increases.

To complement our observational-based flux estimates of dissolved CH₄, we employed three independent atmospheric models to provide top-down constraints of the ocean-atmosphere flux, given the atmospheric concentrations sampled by the aircraft and the ship. This approach also takes potential CH₄ from bubbles into account. We only detected a weak increase of 2 ppb in the atmospheric mixing ratio at the ship location close to bubbles, reflecting the potential enhancement from both dissolved CH₄ and CH₄ from bubbles. We calculated, using a Lagrangian transport model (FLEXPART), the CH_4 enhancements at the ship for all locations that would result from a 1 nmol m⁻² s⁻¹ flux from the area, encompassing the identified and the potential CH₄ seep sites around Svalbard [Sahling et al., 2014] (Figure 2). Running FLEXPART backward in time for all ship positions over the period 20 June to 1 August, the modeled CH₄ enhancement is shown as the yellow line in the supporting information section, Figure S4; compared to the observations, no correlation ($r^2 = 0.003$) is evident. The most sensitive days are the highest 20% modeled peaks (bold yellow line). Using the most sensitive days from this period, we estimate a top-down constraint on the flux from the seep areas of $<2.4\pm1.3$ nmol m⁻² s⁻¹. This estimation assumes that all of the measured 2 ppb variation in the atmosphere is solely due to a flux from the modeled seep areas around Svalbard (Figure 2). Similarly, using a forward chemistry transport model (Oslo CTM3) [Søvde et al., 2012], a flux of 3.8 ± 0.7 nmol m⁻² s⁻¹ was necessary to reproduce the 2 ppb increase in CH₄ at the ship, assuming the same emission region shown in Figure 2. This is equivalent to an annual emission of only 0.06 Tg for a constant flux throughout the year, very small compared to the total global annual emission of ~600 Tg of CH₄ [Kirschke et al., 2013]. In addition, we used the aircraft measurements to provide another independent constrain on the maximum possible CH₄ flux in the region. The aircraft flew transects below 100 m altitude upwind and downwind of the potential seep sites but observed no statistically significant change in CH₄ during these low-level flights; see Figure 1d for altitudes. A Lagrangian mass balance calculation (similar to that employed by O'Shea et al. [2014] leads to an estimated flux of -3.0 ± 17.1 nmol m⁻² s⁻¹. An estimated upper limit on the ocean-to-atmosphere CH₄ flux averaged over the grey shaded area shown in Figure 1d can then be quantified by the mean + 1σ value of 14.1 nmol m⁻² s⁻¹. This represents the maximum possible flux for this area consistent with the aircraft CH₄ measurements and associated uncertainties.

FAAM aircraft measurements were also made in the same location off Prins Karls Forland in a previous Methane in the Arctic Measurement and Modelling (MAMM) campaign in summer 2012 as part of the UK

^bLagrangian particle dispersion model [Stohl et al., 2005; Thompson and Stohl, 2014].

^cChemical transport model [Søvde et al., 2012; Dalsøren et al., 2016].

^dLagrangian mass balance approach [Karion et al., 2013; O'Shea et al., 2014]. Note that the flux constrain based on the flight data is weaker; there was no statistically significant change in downwind CH₄ mixing ratio relative to the measured upwind background, and this is the maximum possible flux that is consistent with the atmospheric flight measurements and associated uncertainties.

Methane in the Arctic Measurement and Modelling (MAMM) project (see Allen et al. [2014] for details). Similarly, any emission from the seep areas was not detectable among the other signals in the aircraft data. Forward calculations, with a different dispersion model, led to very similar conclusions to those of 2014: that an emission flux of a few tens of nmol m⁻² s⁻¹ would have been required to detect the emission in the aircraft data [M. Cain, personal communication, 2016].

In sharp contrast to the flux calculations from the measurement-led approaches discussed here (Table 1), the flux reported by Shakhova et al. [2014] from the East Siberian Arctic Shelf is more than 2 orders of magnitude larger, 70-450 nmol m⁻² s⁻¹ under windy conditions than our measurement-derived maximum for the period. Figure 4e includes a comparison. Part of this large difference can be explained by both higher dissolved CH_4 concentrations in surface waters reported in the Siberian area (up to ~400 nmol L⁻¹) and the higher wind speeds reported by Shakhova et al. [2014]. Table 1 compiles our estimates of the spatially averaged maximal flux in the region, as constrained by the atmospheric observations.

4. Conclusion

Despite the obvious influence of seeps on dissolved CH₄ concentrations in the ocean west of Svalbard in June-July summer 2014, very little CH₄ reaches the atmosphere, neither as bubbles transported nor dissolved gas. The median wind speed was 6.6 m s⁻¹ during our campaign, and the pycnocline remained stable. We suggest that dissolved methane captured below the pycnocline may only be released to the atmosphere when physical processes remove this dynamic barrier. In such a situation, dissolved CH₄ concentrations would rapidly decrease and any large flux would most likely be transient. Consequently, we conclude that large CH₄ releases to the atmosphere with strong impact on the atmospheric levels from subsea sources, including hydrates, do not occur to the west of Svalbard, presently. Shorter periods with large fluxes, particularly during other times of the year such as during ice break-up or storm events, might occur. The role of the pycnocline in this context will be investigated in more detail during long-term ocean observatory recordings in the future.

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