



Diel and seasonal methane dynamics in the shallow and

turbulent Wadden Sea

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- 15 Abstract. The Wadden Sea is a coastal system fringing the land-sea borders of Denmark, Germany, and the 16 Netherlands. The Wadden Sea is extremely productive and influenced by strong variations in physical and
- 17 biological forcing factors that act on time scales of hours to seasons. Productive coastal seas are known to dominate
- 18 the ocean's methane emission to the atmosphere, but knowledge on controls and temporal variations of methane
- 19 dynamics in these vastly dynamic systems are scarce. Here we address this knowledge gap by measuring methane
- 20 inventories and methanotrophic activity at a temporal resolution of two hours over a time period of two days, 21 repeatedly during four successive seasons in the central Dutch Wadden Sea. We found that methane dynamics
- 22 varied between colder and warmer seasons, with generally higher water column methane concentrations and
- 23 methanotrophic activity in the warmer seasons. Efflux of methane to the atmosphere was, on the other hand, lower
- 24
- in the warmer seasons because of lower wind speeds. On a diel scale, tides controlled methanotrophic activity, 25 which increased ~ 40 % at low tide compared to high tide. We estimate that methane oxidizing bacteria reduce the
- methane budget of the Dutch Wadden Sea by only 2 %, while $\sim 1/3$ escapes to the atmosphere and $\sim 2/3$ are flushed 26
- 27 out into the open North Sea at ebb tide. Our findings indicate that tides play a key role in controlling methane
- 28 dynamics and methanotrophic activity and highlight the importance of high resolution and repeated sampling
- 29 strategies to resolve methane dynamics in fast-changing coastal systems.





1 Introduction

1.1 Methane and methane oxidation

Atmospheric methane (CH₄) concentrations have been increasing since industrial times, surpassing 1900 ppb in 2021(Lan et al., 2022) and contributing more than 20 % of total radiative forcing in the atmosphere (Etminan et al., 2016). Due to its relative short atmospheric lifetime of ~10 years (Canadell et al., 2021), reducing methane emissions to the atmosphere could play a key role in global warming mitigation strategies. However, implementation of such strategies requires a thorough understanding of methane sources and sinks. Anthropogenic methane emissions (336 – 376 Tg y⁻¹) are rather well constrained and constitute ~60 % of the total atmospheric budget, but large uncertainties exist around the strength of individual natural sources (Saunois et al., 2020). Especially, methane emissions from marine environments (5 -28 Tg CH₄ y⁻¹) are not well constrained (Weber et al., 2019; Rosentreter et al., 2021).

The inner shelf (0-50 m) water depth) only account for $\sim 3 \%$ of the global ocean surface but are a main source of marine methane emissions to the atmosphere (Weber et al., 2019). In these shallow ecosystems, light availability as well as terrestrial inputs of nutrients support a high diversity of producers and consumers that generate huge quantities of organic matter (Philippart et al., 2009; Beck and Brumsack, 2012). Consequently, rates of organic matter degradation, including methanogenesis in anoxic sediments are high, often leading to elevated levels of free and dissolved methane in sediments and pore waters (Bange et al., 1994; Røy et al., 2008; Wu et al., 2015). Transport of methane-rich porewaters and ebullition of methane bubbles, in return, lead to elevated methane concentrations in the water column (Reeburgh, 2007; Grunwald et al., 2009; James et al., 2016). It is estimated that ~ 5 % of shelf seas surface waters have methane concentrations above 100 nM (Weber et al., 2019). Nevertheless, a substantial amount of dissolved methane is oxidized by aerobic methanotrophic bacteria (MOB), which mediate the aerobic oxidation of methane (MOx) (Reeburgh, 2007):

53
$$CH_4 + 2O_2 \rightarrow CO_2 + H_2O$$
 (1)

Similar to other metabolic processes involving small molecules, MOx discriminates against isotopically heavy methane (i.e. containing ¹³C and ²H (D) instead of ¹²C and ¹H) so that the residual methane pool successively becomes ¹³C and D enriched as a result of ongoing MOx (Barker and Fritz, 1981; Whiticar, 1999).

MOB typically belong to the Gamma- (type I and type X), Alphaproteobacteria (type II), Verrucomicrobia and members of candidate division NC10 (Hanson and Hanson, 1996; Knief, 2015). MOBs build a microbial methane filter in the water column that functions as the ultimate sink for oceanic methane before reaching the atmosphere. Yet, little is known about the controls and capacity of this microbial filter in the inner shelf ecosystems where the vertical distance between the sedimentary source and the atmosphere is short. Factors such as oxygen (Boetius and Wenzhöfer, 2013; Steinle et al., 2017) and methane availability (Mau et al., 2013; James et al., 2016) affect MOx, but also increasing water temperatures play a role by impacting metabolic rates of MOBs (He et al., 2012). The capacity of the microbial MOB filter increases with continuity in the water column (Steinle et al., 2015; James et al., 2016). However, water mass movement induced by temperature changes and wind leads to shifting mixing regimes that disrupt such continuity on a seasonal scale (Gründger et al., 2021). On a daily scale, tides induce currents and currents disrupt continuity and hence can affect MOx, too (Steinle et al., 2015). This disruption of continuity is particularly strong in the extremely dynamic inner-shelf seas where rapid changes in environmental conditions can lead to rapid changes in water column dynamics.

The Wadden Sea, a UNESCO heritage site that consists of the largest continuous tidal flat area worldwide (14.900 km²), is an extremely dynamic system, with major hydrological changes occurring at seasonal to diel time scales. The Wadden Sea stretches for about 500 km along the coast of the Netherlands, Germany and Denmark. Here, we investigated methane dynamics in the Dutch part of the Wadden Sea, that is separated from the North Sea by five barrier islands (Fig. 1). Our aim was to temporally resolve methane dynamics from an hourly to a seasonal scale to determine key controls on methane dynamics and to establish a methane budget for the Dutch Waddensea





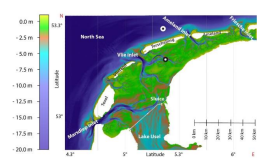


Figure 1. Bathymetry of the western sector of the Dutch Wadden Sea between the Marsdiep and Friesche inlet (modified from Materić et al., 2022). Tidal inlets between barrier islands facilitate water exchange with the open North Sea. The time-series station is located south of the island Terschelling (black mark; 53°19.015 N, 5°22.711 E). The offshore reference station is located 8 km north of Terschelling (white mark; 53°29.190 N, 5°21.449 E).

2 Materials and methods

2.1 Experimental design

A chain of 5 barrier islands (located 5 to 30 km offshore) shelters the Dutch Wadden Sea from waves and strong westerly winds. Between these barrier islands, large volumes of water are transported in and out the Dutch Wadden Sea with the rhythm of the tides through deep tidal inlets, such as the Marsdiep (most western point of the Dutch Wadden Sea) and the Vlie inlet (Duran-Matute et al., 2014). Our fixed mooring station (53°19.015 N, 5°22.711 E) is in a branch of the Vlie inlet between the island of Terschelling and the mainland, roughly in the middle of the Dutch Wadden Sea (Fig. 1). This location was chosen as it remains submerged at low tide and lays in-between the Wadden Sea's landward and offshore termination. The water flowing by this station thus equally integrates the tidal flat area, mostly during ebb tide, as well as the inflowing North Sea water during high tide. Also, the station was relatively far away from the port of Harlingen (~ 20 km) so that a potential influence of methane rich port waters is minimized. The reference station was located 8 km north of the island Terschelling in the North Sea (53°29.190 N, 5°21.449 E).

Samples were recovered with the R/V Navicula during 4 sampling campaigns, respectively in winter (19 February 2019 – 21 February 2019), spring (23 April 2019 – 25 April 2019), summer (22 July 2019 – 24 July 2019) and autumn (11 November 2019 – 13 November 2019). During each campaign, we conducted hourly CTD casts with discrete water sampling over a two-day period. During CTD casts, water mass properties (temperature, salinity, density) and oxygen concentrations were measured continuously using a Sea-Bird (SBE911) + conductivity–temperature–depth (CTD) system. Discrete water samples were recovered with Niskin bottles from 1 and 3 m water depth and, upon recovery, immediately sampled for subsequent analyses of water column constituents (methane concentrations, methane isotopic composition and methane oxidation rates).

Sediment samples were retrieved using a boxcorer, and upon recovery, subsampled with small pushcores (diameter 7 cm, \sim 18 cm sediment recovery). Pushcores were subsampled for methane concentrations by taking every 2 cm 5 mL of sediment that was quickly added to 60 mL glass bottles containing 30 mL of a saturated NaCl brine solution and the bottles were imidiately sealed with butyl rubber stoppers. Atmospheric flask samples (250 ml) were taken hourly at \sim 10 m above the sea surface, in winter and spring. In summer and autumn, atmospheric methane concentrations were continuously measured using a cavity ringdown spectrometer (CRDS, Picarro model G2301).

2.2 Dissolved methane concentrations and stable isotope ratios

Dissolved methane concentrations were determined using a headspace (HS) technique (Green, 2005). In brief, immediately upon CTD recovery, 260 mL glass serum bottles were filled HS-free, closed with black-butyl rubber stoppers (Rubber B.V. the Netherlands) and crimp-top sealed. Next, we added a 5 mL N₂ headspace and fixed the sample with 5 mL NaOH solution (25 % w/v). HS methane concentrations of sediments and dissolved methane were measured in our home laboratories with a gas chromatograph (GC; Thermo Scientific FOCUS GC equipped with a Restek stainless steel column HS-Q 80/100 SS GEN config (length 2 m, 2 mm ID, 1/8 OD) with flame

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- 117 ionization detection). The instrument was calibrated with a certified 100 ppm methane standard (Scott Specialty
- 118 Gases Netherlands B.V.).
- 119 Similarly, seawater aliquots were taken for methane stable carbon and hydrogen isotope measurements, but these
- 120 samples were fixed with 60 µl HgCl₂ (2 mM). A continuous flow isotope ratio mass spectrometry (CF-IRMS)
- 121 system was used to quantify D-CH₄ in the gas phase (Thermo Delta Plus XL, Thermo Fisher Scientific Inc.,
- 122 Germany) as described previously (Röckmann et al., 2016; Jacques et al., 2021). Isotopic values are represented
- in the delta notation against the international reference standard VSMOW (δD). To monitor precision and 123
- 124 accuracy, sample measurements were alternated with measurements of an inhouse air standard (cross calibrated
- 125 against certified reference standards) containing 1975.5 ppb methane with a δD -90.81 \pm 1.1 \%. We constructed a
- 126 two-endmember mixing model (Mariotti et al., 1981; Jacques et al., 2021) and a Rayleigh fractionation model.
- 127 This was done to investigate whether enrichment of D in the residual methane was caused by MOx, which is
- 128 known to discriminate against heavy isotopes (Barker and Fritz, 1981; Whiticar, 1999), or by mixing with
- 129 comparably heavy atmospheric methane (see supplementary methods).

130 131

2.3 Methane oxidation rate measurements

- 132 MOx was determined by ex-situ incubations with trace amounts of ³H-labelled methane as described previously
- 133 (Niemann et al., 2015). Briefly, aliquots from each Niskin bottle were filled HS-free in 20 mL glass vials in
- 134 triplicate, sealed with grey-bromobutyl stoppers that are known to not hamper methanotrophic activity and
- 135 amended with 5 μL of ³H-CH₄/N₂ (4.5 kBq, American Radiolabeled Chemicals, USA). Samples were incubated
- 136 for 72 h in the dark at in situ temperature. Activities of residual C³H₄ and the MOx product ³H₂O were measured
- 137 by liquid scintillation counting.
- 138 MOx first-order rate constant (k) was determined from the fractional tracer turnover (Reeburgh, 2007):

139
$$k = \frac{{}^{3}\text{H}_{2}\text{O}}{{}^{3}\text{H}_{2}\text{O} + \text{C}^{3}\text{H}_{4}} \times \frac{1}{\text{t}}$$
 (2)

140

- 141 where t is incubation time in days. k was corrected for (negligible) tracer turnover in killed controls (KC, fixed
- 142 with 100 µl HgCl₂ directly after sampling) and multiplied with dissolved methane concentrations [CH₄], yielding
- 143 MOx:

144
$$MOx = (k - k_{KC}) \times [CH_4]$$
 (3)

145

146 Diffusive fluxes of methane

- The diffusive sea-air methane flux was calculated based on a boundary layer model that consider the relation 147
- 148 between wind, temperature and methane concentrations in the atmosphere and a well-mixed surface water layer
- 149 (Wanninkhof, 2014):

150
$$F = (pCH_{4w} - pCH_{4a}) K_0 k_{CH_A}$$
 (4)

- 151 F denotes the diffusive methane flux, pCH_{4a} and pCH_{4w} (in atm) are the partial pressures of methane in the air and
- 152 in the well-mixed surface water layer, respectively. pCH_{4a} was measured with a Picarro G2301 gas concentration
- 153 analyser on board. pCH_{4w} was determined from surface water methane concentrations (see above). K₀ is the 154 methane solubility in mol m⁻³ atm⁻¹ (Wiesenberg and Guinasso, 1979) and was calculated from temperature and
- 155 salinity obtained from corresponding CTD casts. k_{ch4} is the methane gas transfer velocity in m d⁻¹ which was 156 calculated using wind speed (U), the Schmidt number (Sc_{CH4}) and the normalised gas transfer velocity (k_{660})
- 157 according to (Wanninkhof, 2014):

158
$$k_{\text{CH}_4} = 0.251 U^2 \left(\frac{\text{Se}_{\text{CH}_4}}{660}\right)^{-0.5}$$
 (5)

- 159 Wind speed was measured on board at 10 m above sea level. The Schmidt number describes the ratio between
- 160 kinematic viscosity of water and the gas diffusion coefficient, which relates the different k-values for different
- 161 gases (Jähne et al., 1987; Wanninkhof, 2014).





2.5 Statistical analysis

A principial component analysis (PCA) was carried out to study the relationship between environmental variables and methanotrophic activity. The input variables for the PCA were temperature, salinity, density, *k*, MOx, and dissolved methane concentrations. Prior to running the PCA, the variables were centered and scaled. We utilized the R software and the 'FactoMineR' package (Lê et al., 2008) for the PCA analyses.

3 Results

3.1 Dynamics of sea water properties

Water column temperature varied between seasons and ranged from 6.3 °C to 24 °C (Fig. 2, Table 1). A clear distinction could be made between colder seasons (autumn and winter) in which temperature ranged from 6.3 °C to 9.1 °C and warmer (spring and summer) seasons where temperatures ranged from 14.2 °C to 24 °C. Water temperatures at the reference station were similar in winter (6.9 °C) but colder in spring (10.5 °C) and summer (20.3 °C) and warmer in autumn (11.8 °C) when compared to the Wadden Sea.

Table 1. Average seawater temperature, salinity, and density at the time-series station (central Dutch Wadden Sea) and
 reference station (offshore Terschelling, North Sea)

	Autumn	Winter	Spring	Summer
Temperature (°C)	8.4 ± 0.4	6.7 ± 0.2	15.0 ± 0.6	22.1 ± 0.8
Salinity (psu)	22.4 ± 1.9	23.4 ± 1.8	31.3 ± 0.2	30.6 ± 0.6
Density (σ _t)	17.4 ± 1.4	18.3 ± 1.4	23.1 ± 0.2	20.8 ± 0.6
	Autumn Ref st.	Winter Ref st.	Spring Ref st.	Summer Ref st.
Temperature (°C)	11.8 ± 0.01	6.8 ± 0.1	10.4 ± 0.12	20.0 ± 0.1
Salinity (psu)	31.3 ± 0.01	32.2 ± 0.1	31.8 ± 0.03	32.3 ± 0.03
Density (σ _t)				

On a diel scale, variation in water temperature were related to the tidal phase. In winter, spring, and summer, maximum water temperatures were observed around low tide (LT, here defined as the time when we encountered the lowest water depth during CTD casts, Fig. 2). This was 7.2 °C in winter, 17.3 °C in spring and 24.1 °C in summer. Minimum water temperatures were around high tide (HT, high tide, here defined as the time when we encountered maximum water depth during CTD casts, Fig. 2). This was 6.3 °C winter, 14.2 °C in spring, and 20.9 °C in summer. In autumn, this pattern was inverse with minimum water temperatures at LT (7.6 °C) and maximum at HT (9.1 °C).

Like temperature, salinity differed strongly between colder (18 - 27 psu) and warmer seasons (29 - 32 psu; Fig. 2, Table 1). Furthermore, salinity was higher during HT irrespective of season. Salinity levels at the reference station in the North Sea were stable (31.3 – 32.3 psu) without obvious diel and seasonal fluctuations. Changes in density were caused by salinity rather than temperature during all four seasons, with one exception in spring: after 28 hours of the time series, salinity remained stable, but water temperatures decreased which lowered water density.





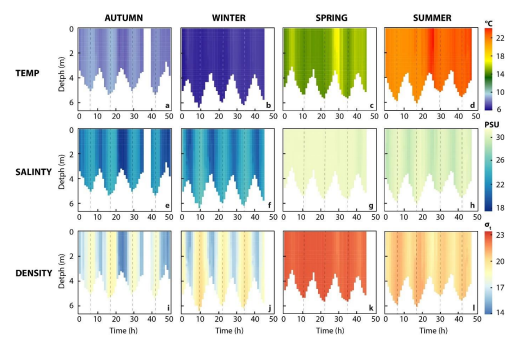


Figure 2. Properties of sea water. (a-d) Spatiotemporal distribution of temperature, (e-h) salinity and (i-l) density. Dashed line indicates high tide.

3.2 Methane dynamics

3.2.1 Methane concentrations in the water column and in sediments

Water column methane concentrations showed a high degree of variability and were clearly distinguishable between the colder and warmer seasons (Fig. 3A-D, Table 2). We found a significant difference in average methane concentrations between 1 m (16.0 nM) and 3 m (17.6 nM) water depth in winter (p \leq 0.007, Welch's t-test). In autumn, methane concentrations were also lower at 1 m (15.5 nM) than at 3 m (16.2 nM) water depth, but the difference was not significant. However, it is noteworthy that the methane concentrations at the beginning of the time-series were around 35 nM and rapidly decreased to values below 15 nM within one day. During warmer seasons, average methane concentrations were similar at the surface and in deeper waters, i.e. 40.9 nM (1 m) and 41.3 nM (3 m) in spring and 69.2 nM (1 m) and 69.4 (3 m) in summer. Methane concentrations at our reference station were \sim 3 nM in winter, spring, and autumn and \sim 6 nM in summer and thus far lower when compared to the Wadden Sea.

On a diel scale, methane concentrations varied during all seasons, roughly matching the tidal regime. In spring at LT, depth-averaged methane concentrations were $42.6\,\mathrm{nM}$, but decreased by $\sim25\,\%$ to $34.2\,\mathrm{nM}$ at HT. This pattern also occurred in autumn where methane concentrations decreased by $21\,\%$ from $17.4\,\mathrm{nM}$ at LT to $14.4\,\mathrm{nM}$ at HT. In winter ($14.7\,\mathrm{nM}$ at LT and $14.3\,\mathrm{nM}$ at HT) and summer ($72.5\,\mathrm{nM}$ at LT and $71.3\,\mathrm{nM}$ at HT), the difference between LT and HT was smaller (Table 2).

Sediment methane concentrations increased with depth during all seasons (Fig. S1 in the Supplement). Concentrations were similar in autumn (0.5 – 2.2 μ M), winter (0.4 - 0.6 μ M) and spring (0.5 - 0.9 μ M) but in summer, we found highly elevated sediment methane concentrations ranging from 3.6 to 18.7 μ M. The high concentrations in sediments during the summer season are in line with an increase in dissolved methane concentrations in the water column.





Table 2. Methane dynamics in the Dutch Wadden Sea. Average and standard deviation of methane concentrations, k, MOx and δ D-CH₄ during four seasons in 2019. Values represent averages for 1 and 3 m water depth (averaged over the two-day time series recorded for each season) as well as for low and high tide only (averaged over depth). LT = minimal water depth during CTD casts, HT = maximum water depth during CTD cast. Average wind speed and methane efflux to the atmosphere are averaged over the two-day time series recorded for each season. ns = not sampled.

	Autumn	Winter	Spring	Summer			
Methane concentration (nM)							
1 m water depth	15.5 ± 5.8	16.0 ± 2.4	40.9 ± 9.2	69.2 ± 21.4			
3 m water depth	16.2 ± 5.7	17.6 ± 3.0	41.3 ± 8.9	69.4 ± 22.4			
Low tide	17.4 ± 9.7	14.7 ± 2.1	42.6 ± 6.9	72.5 ± 36.1			
High tide	14.4 ± 1.6	14.3 ± 0.6	34.2 ± 10.7	71.3 ± 27.4			
Reference station	3.3	3.1	3.7	6.6			
k (d·1)							
1 m water depth	0.03 ± 0.01	0.02 ± 0.01	0.03 ± 0.01	0.07 ± 0.02			
3 m water depth	0.03 ± 0.02	0.03 ± 0.01	0.03 ± 0.02	0.06 ± 0.02			
Low tide	0.05 ± 0.01	0.03 ± 0.01	0.05 ± 0.01	0.08 ± 0.02			
High tide	0.03 ± 0.01	0.03 ± 0.02	0.02 ± 0.01	0.06 ± 0.02			
Reference station	0.01	0.0004	0.02	0.04			
MOx (nM d ⁻¹)							
1 m water depth	0.48 ± 0.22	0.39 ± 0.21	1.16 ± 0.61	4.41 ± 1.49			
3 m water depth	0.54 ± 0.34	0.52 ± 0.27	1.33 ± 0.71	4.33 ± 1.84			
Low tide	1.05 ± 0.48	0.47 ± 0.24	2.02 ± 0.42	5.24 ± 2.33			
High tide	0.50 ± 0.16	0.43 ± 0.31	0.59 ± 0.19	4.23 ± 2.13			
Reference station	0.03	0.001	0.07	0.23			
δD-CH ₄ (‰)							
1 m water depth	-219 ± 31	ns	ns	-250 ± 17			
3 m water depth	-224 ± 27	ns	ns	-250 ± 14			
Low tide	-208 ± 41	ns	ns	-227 ± 1			
High tide	-227 ± 13	ns	ns	-265 ± 3			
Methane sea-air flux (μmol m ⁻² d ⁻¹)							
Wind speed (m s ⁻¹)	8.0 ± 2.1	8.3 ± 1.4	7.9 ± 2.7	3.8 ± 1.6			
Methane flux	40.2 ± 28.1	38.7 ± 14	144.8 ± 98	72.9 ± 52			
Atmosphere conc. (ppm)	2.0 ± 0.03	2.12 ± 0.19	2.02 ± 0.15	2.14 ± 0.15			





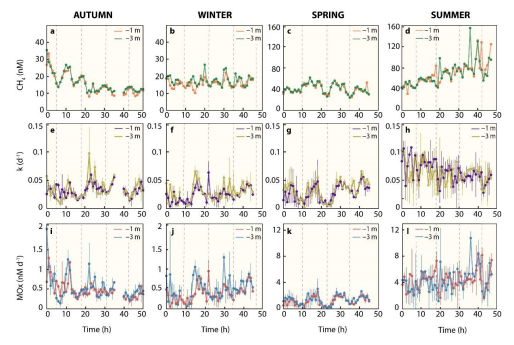


Figure 3. Methane dynamics. (a-d) Dissolved methane concentration, (e-d) first-order rate constant, (i-l) methane oxidation rates. Note that for dissolved methane concentrations in colder seasons, autumn and winter, the y-axis differs from warmer seasons, spring, and summer. Dashed line indicates high tide.

3.2.2 Methane oxidation rates

Similar to methane concentrations, we observed strong seasonal differences in MOx (Fig. 31-L, Table 2). Depth-averaged MOx in spring (1.2 nM d^{-1}) and summer (4.4 nM d^{-1}) was ~ 3 and ~ 9 – fold higher than in winter (0.5 nM d^{-1}) and autumn (0.5 nM d^{-1}). MOx at 1 m and 3 m water depth statistically differed from each other in winter (p \leq 0.01, Welch's t-test), but not in spring, summer, and autumn. MOx at the reference station was < 5 % of MOx in the Wadden Sea, with maxima found in summer (0.2 nM d^{-1}).

On a diel scale, MOx showed fluctuations during all seasons. In general, depth-averaged MOx was higher during LT compared to HT. In autumn average MOx at LT (0.79 nM d^{-1}) was about 2 - fold higher and significant different from MOx at HT $(0.38 \text{ nM d}^{-1}, p \le 0.03, \text{Welch's t-test})$.

In winter, the difference between MOx at LT (0.47 nM d⁻¹) and HT (0.43 nM d⁻¹) was small. In spring, depth-averaged MOx at LT (2.02 nM d⁻¹) was about 4 – fold and significantly ($p \le 6.4 \times 10^{-6}$, Welch's t-test) higher than during HT (0.58 nM d⁻¹). In summer, MOx was high at both, LT (5.2 nM d⁻¹) and HT (5.4 nM d⁻¹). Similarly, k was substantial higher (16 – 63 %) at LT than HT in all seasons (Fig. 3E-H, Table 2). In fact, the difference in depth-averaged k between LT and HT was significant in autumn ($p \le 0.003$, Welch's t-test) and spring ($p \le 6 \times 10^{-5}$, Welch's t-test, Table S1 in the Supplement).

3.2.3 Stable hydrogen isotope signatures

The stable hydrogen isotope composition of dissolved methane was only measured in autumn and summer (Fig. 4, Table 2). In autumn, average δD -CH₄ over the entire time-series was -219 % at 1 m water depth and -224 % at 3 m water depth, but there was a generally strong trend towards higher δD -CH₄ values over the two-day period from about -260 % to about -180 %. In summer the mean δD -CH₄ values were homogenous throughout the water column (-250 %) and generally lower than in autumn. Except for the first full tidal cycle in autumn, the results showed a tidal imprint on δD -CH₄ values with higher δD -CH₄ values at LT and lower values at HT independent of depth and season (Fig. 4).



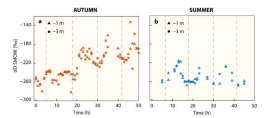


Figure 4. Progression of δD -CH₄ signatures in (a) autumn and (b) summer at 1 m and 3 m water depth. Vertical dashed line indicates high tide.

In addition to tidal patterns, the δD -CH₄ values in autumn were substantially higher at lower methane concentrations (< 21 nM, Fig. 5). Linear mixing alone of (i) well-mixed surface waters in equilibrium with atmospheric methane and (ii) the maximum methane concentration in the water column, both concentrations with their associated isotopic signatures, would result in concentration/isotope data as depicted by the mixing lines in Fig. 5. Results in autumn clearly deviated from this mixing line at low methane concentrations. On the other hand, the open system Rayleigh fractionation model that we ran for low methane concentration in autumn yielded an ϵ -value of -97 ‰ and matched the steep rise in δD -CH₄ with decreasing methane concentration much better. This directly indicates that MOx is the dominant mechanism driving δD -CH₄ to higher values at low concentrations.

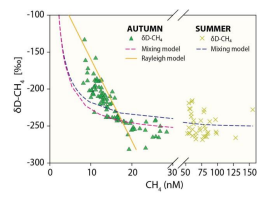


Figure 5. Methane concentration versus δD -CH₄ - mixing and oxidative removal in autumn and summer. Dashed lines show methane concentration/isotope dynamics determined with a two-endmember mixing model considering (i) well-mixed Wadden Sea surface waters and (ii) methane charged waters as endmembers. Methane concentration and stable hydrogen isotope composition following oxidative removal according to a Rayleigh model for low methane concentrations are depicted as a solid line. Samples with methane concentrations < 21 nM (δD -CH₄ = ~ -217 %) in autumn and < 61 nM (δD -CH₄ = ~ -244 %) in summer were defined as the methane source signal and thus starting point of the Rayleigh fractionation model. The apparent isotope enrichment (ε see also Fig. S2 in the Supplement) was -97 % in autumn. Neither the mixing nor the Rayleigh model are well constrained for δD -CH₄ in summer; the mixing line is thus only shown for comparison and ε could not be calculated.

3.2.4 Diffusive efflux to the atmosphere

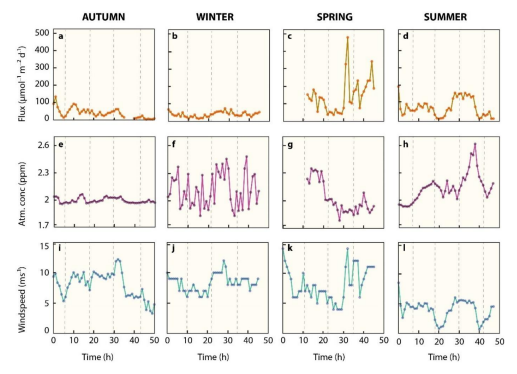
The water column in the Wadden Sea was consistently methane supersaturated (> 8 nM) with respect to atmospheric equilibrium (~ 2.6 nM) during all sampling campaigns (Fig. 3, Table 2), which indicates a continuous release of methane from the water to the atmosphere throughout the measurement series. Atmospheric concentrations were similar ranging from 1.8 to 2.6 ppm, with relatively constant concentrations in autumn and more erratic concentrations in winter, spring, and summer (Fig. 6e-h, Table 2). Noteworthy is the sharp increase of atmospheric methane from 2 to 2.6 ppm between 29 and 38 hours in summer before decreasing again to 2 ppm.

Windspeeds in autumn, winter, and spring where relatively high (typically $> 5 \text{ ms}^{-1}$) when compared to calmer conditions in summertime (typically $< 5 \text{ ms}^{-1}$, Fig. 6i-1, Table 2). As a result of the strong but variable wind forcing, diffusive methane fluxes fluctuated in magnitude within, and between season (Fig. 6a-d, Table 2). Average diffusive fluxes in autumn and winter were with $< 40 \mu$ mol m⁻² d⁻¹ about 4 – fold lower than in spring and 2 – fold lower than in summer. Maximum efflux (479 μ mol m⁻² d⁻¹) in spring occurred after the wind velocity increased





282 rapidly from 6 m s $^{-1}$ to 14 m s $^{-1}$ within two hours and methane concentrations slightly increased from 38 nM to 45 nM.



 $\label{eq:Figure 6. Diffusive methane flux. (a-d) Sea surface atmosphere methane fluxes, (e-h) Seasonal atmospheric methane concentrations, (i-l) local wind speed. Vertical dashed lines indicate high tide.$

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3.3 Statistical Analysis

To study the relationship between environmental variables and methanotrophic activity, we conducted a Principal Component Analysis (PCA). The outcome explained 92 % of the data variability on the first two components (Fig. 7, Table S2 in the Supplements). The main gradient (PC1: 69 %) showed a contrast between autumn/winter and summer and spring. Temperature, salinity, methane concentrations and MOx peaked in summer and spring, while lower values were measured in winter/autumn. The relatively small ellipse in spring indicates that samples show more similarity than in other seasons. The second gradient distinguished the summer from the spring samples based on higher *k* and density in the latter (PC2: 23 %).

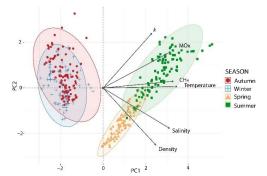


Figure 7. Principal Component Analysis (PCA) of environmental conditions across seasons in the Dutch Wadden Sea. Biplot of a PCA of the explanatory variables as vectors (in black) and observations (marks) of each season on the first (x-axis, PC1) and second principal component (y-axis, PC2). Coloured concentration ellipses (size determined by a 0.95-probability level) show the observations grouped by season. The magnitude of the vectors (line length) shows the strength of their contribution to the PCs. Vectors pointing in similar directions indicate positively correlated variables and vectors at angles > 90° indicate no correlation.

4 Discussion

The Wadden Sea is a highly productive ecosystem (Philippart et al., 2009) where the decay of organic matter supports high rates of methanogenesis in sediments (Røy et al., 2008; Wu et al., 2015), Which in return leads to high methane concentrations in the Wadden Sea's water column (Grunwald et al., 2007; Grunwald et al., 2009). Little knowledge, however, exists on the variability of methane dynamics on short time scales of hours to days or between seasons and the underlying controls on this variability. Here, we measured water column methane concentrations, methane oxidation and the oceanographic regime as well as atmospheric methane mixing ratios and wind velocity in the Dutch sector of the Wadden Sea for two days during four consecutive seasons in 2019.

4.1 Water column properties

In general, we found a clear distinction between colder (autumn and winter) and warmer (spring and summer) seasons (Figs. 2, 7). North Sea waters with incoming tide, flow through tidal inlets that in turn branch into successively smaller tidal creeks in which water-flow direction alternates with the tidal phase. This then led to increasing water temperatures in autumn but decreasing water temperatures in winter. The high temperature of Wadden Sea waters during incoming tide in autumn can be explained by the fact that the shallow Wadden Sea cools rapidly once the summer is over, while the North Sea's large water volume takes longer to cool down.

Salinity levels were on average lower in colder seasons compared to warmer seasons, likely because land runoff and ground water discharge are typically higher in autumn and winter because of the overall higher precipitation levels during the cold seasons (Van Aken, 2008). A higher level of freshwater inflow from land was also evident from the rapidly dropping salinity levels during falling and LT in autumn and winter (Fig. 2). This freshening effect is amplified at times when the Dutch Ministry of Infrastructure and Water management (Rijkswaterstaat) opens water gates to discharge excess water from lake IJsel (Fig. 1), which occurs more often in colder seasons due to increased input of precipitation, groundwater discharge as well as surface and riverine discharge to the lake. During the warmer and dryer seasons, water gates are mostly kept closed to ensure that the lake's water level stays high. However, freshwater inflow into the Wadden Sea was evident during all seasons because incoming North Sea water generally increased salinity levels at HT independent of sampling time.





4.2 Differences in methane concentrations and isotopic signatures on time scales of seasons

Sediment and water column methane concentrations were highly elevated in summer (Figs. 3, S1 in the Supplement and Table 2). In fact, average sediment methane concentrations increased 17 - fold in summer compared to spring two months earlier. This increase is probably related to the remineralization of the spring phytoplankton bloom that takes place in the months of April and May (Philippart et al., 2009) leading to elevated rates of methanogenesis in anaerobic sediments (Beck and Brumsack, 2012). A time lag of one to two months between the peak of the spring bloom and methane release from sediments was also observed in the Baltic Sea (Bange et al., 2010). In addition to elevated inputs of organic matter, increasing water temperatures towards summer further enhances microbial methanogenesis in sediments (Yvon-Durocher et al., 2014). We indeed observed lower methane concentrations in autumn and winter compared to spring and summer, which is most likely related to both reduced organic matter input and colder temperatures. It has to be noted that the sediment methane concentrations presented here are comparably low as sediment methane concentrations close to saturation levels were previously found at other locations in Wadden Sea sediments (Røy et al., 2008; Wu et al., 2015). We did not measure sulphate concentrations, but the methane profiles indicate that we only reached the upper part of the methane-sulphate transition zone below of which methanogenesis proceeds. Also, sediment methane concentrations can be variable on spatial scales of metres. Depending on the hydrographic regime, the methanesulphate transition zone can be metres below the tidal flat sediments (Wu et al., 2015), but pore water flow can also transport reduced compounds such as sulphide and methane to the sediment surface (Røy et al., 2008).

Methane release from sediments and the relatively low wind speed (and thus relatively low forcing to drive diffusive efflux) in summer led to charging of the water column with methane. MOx discriminates against isotopically heavy methane and thus causes an isotopic enrichment of residual methane. Thes isotopic discrimination effect manifests more pronouncedly at low methane concentrations. Indeed, we found more pronounced MOx induced isotopic discrimination effects in autumn at low methane concentrations (< 21 nM). At higher methane concentrations (> 21 nM) values were more depleted and were comparable to summer δ D-CH₄ values. We relate the δ D-CH₄ values (< -217 % in autumn and < -244 % in summer) at higher methane concentrations (> 21 nM in autumn and > 61 nM in summer) to the δ D-CH₄ source signal (Fig. 4A, 5). At these concentrations, the isotope effect imposed by MOx is masked by the high background methane and/or is overprinted by methane entering the water column from sediments.

4.3 Differences in MOx on seasonal time scales

The activity of MOBs in the water column is determined by the availability of methane oxygen, nutrients, and the size of the standing stock of the MOB community (Reeburgh, 2007; Crespo-Medina et al., 2014; Steinle et al., 2015). The Wadden Sea water column is a nutrient rich and typically oxygenated environment, we hence argue that nutrient and O₂ availability are not a limiting factor for MOB activity. However, MOBs in the Wadden Sea need to cope with high fluctuations in temperature, salinity, and methane availability (see above). We did not measure the size of the MOB community; nevertheless, it seems likely that the highly variable water column properties with admixture of different water masses and resuspension of particles effects the standing stock of the MOB community and/or its activity.

Previous studies showed that elevated salinity often led to an immediate decrease in MOx in terrestrial/lacustrine systems (Ho et al., 2018; Zhang et al., 2023). Likewise, marine methanotrophs seem to function best at salinity levels of > 20 psu (Osudar et al., 2017), while a sudden decrease in salinity can strongly inhibit MOx (Hirayama et al., 2013; Tavormina et al., 2015). This begs the question if waters, with rapidly changing salinity levels such as the Wadden Sea, are environments that are rather not conducive for MOx, in particular in colder months where salinity levels may drop to ~ 20 psu because of elevate freshwater influx (see above). While MOx was indeed lower in autumn and winter, the relative decrease in MOx was moderate in comparison to the previous literature findings (Osudar et al., 2017; Zhang et al., 2023). Also, autumn and winter are colder and defined by lower methane levels, which likely reduces MOx further. Across seasons, the PCA (Fig. 7), and Pearson correlation coefficients of pairs of variables (Fig. S3 in the Supplement) indicated that MOx (or *k*) and salinity (or density) are not or only weakly correlated. The Wadden Sea thus seems to host a euryhaline MOB community that contrast with MOB communities from terrestrial/lacustrine (Zhang et al., 2023) and oceanic origin (Osudar et al., 2015), which seem less able to cope with varying salinity levels.

Sediments and the water column in the Wadden Sea are increasingly fuelled with methane when ambient temperatures rise. The higher availability of methane could then enhance methanotrophic activity (Reeburgh, 380 2007). Indeed, we found a seasonal imprint with highest MOx levels in summer that were 3-fold higher than those





observed in spring, 9-fold higher than in autumn and 10-fold higher than in winter (Table 2). A correlation between methane, temperature and MOx was also apparent from the PCA (Figs. 7, S3 in the Supplement). We note that not only MOx, but also the first order rate constant k was stimulated by higher methane concentrations (MOx is a function of k and [CH₄], see eq. 3) and temperature. A positive effect of methane on MOx and k is often associated with changes in methane concentrations over several orders of magnitude (Crespo-Medina et al., 2014; James et al., 2016). Here we found that k doubled in summer compared to spring, while methane concentrations were only 30 nM higher, i.e., 1.7-fold. This suggests that a combination of methane availability and temperature determined k in our study. I.e., the MOBs may have been stimulated on the enzymatic level. However, the fact that k remained stable in colder seasons with low water temperatures, suggest that additional factors, likely MOB community size (Steinle et al., 2015), might play a more important role in maintaining k. For example, MOBs from sediments can be resuspended into the water column due to tidal currents, or transported from sediments to the water-column with bubbles as has been found at other cold seeps (Steinle et al., 2016; Jordan et al., 2020; Jordan et al., 2021). Resuspension could thus be a key driver of the Wadden Sea water column MOB communities, with major consequences for maintaining a microbial filter under less favourable conditions.

4.4 Methane dynamics on times scales of hours to days.

Strong hydraulic dynamics are an important characteristic of the Dutch Wadden Sea, with tidal currents interchanging a large water volume with the North Sea twice per day (Gräwe et al., 2016). With the change in tidal phase, the hydrostatic pressure changes rapidly with water depth, which triggers porewater flow (tidal pumping, (Røy et al., 2008)) but may also trigger expansion and ebullition of gas bubbles (Schmale et al., 2015; Jordan et al., 2020). Similar effects are caused by tidal currents flowing over bathymetric features, which triggers pore water flow, too, and additionally resuspends sediments and MOBs into the water column (Bussmann, 2005; Abril et al., 2007; Røy et al., 2008). On the other hand, incoming water from the open North Sea contains relatively low amounts of methane (<6 nM as measured at our reference station), hence, this will dilute the Dutch Wadden Sea's methane content, and outflowing water will export methane from the Dutch Wadden Sea main water body.

Temporal patterns of methane concentration and MOx indeed correlated well with tidal oscillation (Figs. 3, S1 in the Supplements, Table 2). Independent of the seasons, methane concentrations and MOx were elevated at LT. The tidal effect seemed most pronounced in spring where at LT, methane concentrations (1.3 – fold), *k* (2.5 – fold) and MOx (4 – fold) were higher than at HT, independent of depth. We found it surprising that, just like methane concentrations, *k* also was substantially higher during LT compared to HT independent of seasons and despite an overall lower salinity at low tide (Figs. 3, S1 in the Supplement, Table 2). To the best of our knowledge, this has not been described before. It seems unlikely that the MOB community substantially grew or that the velocity of the MOB's metabolism increased/decreased in a time frame of a few hours. We rather argue that the observed oscillation is caused by a likewise oscillation of shear force and hydrostatic pressure, leading to resuspension of MOBs from sediments as well as elevated release of methane form the sea floor.

Grunwald et al. (2007 and 2009) conducted time-series measurements in the German sector of the Wadden Sea near the island of Spiekeroog. There, absolute methane concentrations were ~ 3 – fold higher in spring and summer and ~ 15 – fold higher in winter when compared to our study. This might be related to local factors, for example the vicinity of the estuaries of the rivers Eems and more importantly Weser close by Spiekeroog, which increase the background methane concentrations in this sector of the Wadden Sea. Like in our study, Grunwald et al. (2007, 2009) also reported on a strong influence of tides, with highest methane concentrations at low tide, probably related to tidal pumping, while inflowing waters showed concentrations typical for the open North Sea in the German Bight. The temporal aspects and processes determining methane dynamics discussed in our work are thus not a local feature but applicable to the entire Wadden Sea and likely to other mud flat areas influenced by tides, too.

4.5 Methane emissions from the Dutch Wadden Sea

Surface waters were supersaturated in methane with respect to the atmospheric equilibrium during all seasons; the Wadden Sea is consequently a constant source of methane to the atmosphere. Just as for dissolved methane concentrations and MOx, methane efflux to the atmosphere was higher during warmer seasons compared to colder seasons. This was primarily driven by methane concentrations rather than wind velocity: wind speeds were similar in autumn, winter, and spring, but 2–fold higher methane concentrations in spring translate to a 4–fold higher sea-air flux when compared to autumn and winter. In summer, meteorological conditions were dominated by a heat wave with extremely low wind speeds. This resulted in a comparably low methane efflux to the atmosphere (though still higher than during the colder seasons) leading to an accumulation of methane in the water column. Previously described diffusive methane fluxes at coastal systems vary over several orders of magnitude and appear



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site specific. At the Baltic sea coast, fluxes of up to 15 μ mol m⁻² d⁻¹ were reported (Bange et al., 2010; Steinle et al., 2017), while in arctic shelf seas, diffusive fluxes of up to 240 μ mol m⁻² d⁻¹ were found (Thornton et al., 2016). In comparison, estuarine research along the European Atlantic coast found a median flux of 130 μ mol m⁻² d⁻¹ (Middelburg et al., 2002), which is comparable to the fluxes that we found in the Dutch Wadden Sea (~39 to 145 μ mol m⁻² d⁻¹).

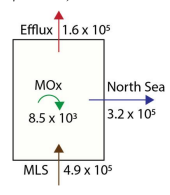


Figure 8. Methane budget for the Dutch Wadden Sea is calculated based on values for the Wadden Sea's geometry, tidal displacement volume, and biogeochemical parameters as discussed in the text. MLS stands for methane liberation from sediments. All values are presented as mol CH₄ d⁻¹.

Towards a roughly estimated methane budget for Dutch Wadden Sea, we combined our diffusive flux, MOx and methane concentration data (Fig 8.) as well as estimates of the Wadden Sea water volume and tidal prism. Our flux estimates (Table 2) translate to an annual average sea surface-atmosphere flux of 74 µmol m2 d-1. Extrapolating this to the area of Dutch sector of the Wadden Sea (~2200 km²; (Materić et al., 2022) implies that 1.6×10^5 mol CH₄ d⁻¹ escapes from the Dutch Wadden Sea to the atmosphere (Table 2). The average water volume of the Dutch Wadden Sea is about 5.15 km³ (Materić et al., 2022); hence the annual average of 1.7 nM d⁻¹ of MOx translates to 0.09×10^5 mol CH₄ d⁻¹ that is oxidized in the water column by MOBs. In addition to atmospheric efflux and microbial consumption, methane rich waters are also flushed into the North Sea. To estimate this, we simplified that the total tidal prism of 4.5 km³ (Gräwe et al., 2016) is an approximation of the net amount of water that leaves the Wadden Sea during LT. With respect to our measured mean methane concentration (36.8 nM), about 1.6×10^5 mol of methane are thus flushed towards the North Sea twice daily, i.e., 3.2×10^5 mol per day. A large uncertainty in this calculation is caused by the delay of ~ 3 hours in tidal phases between the Western and Eastern part of the Dutch Wadden Sea. In other words, methane-rich waters are flowing out of the tidal inlet in the West can be entrained in the current that starts flowing back into the Wadden Sea at eastern tidal inlets. Therefore, the net loss of methane to the Wadden Sea is probably lower than described above. Still, data from our reference station show only slightly oversaturated methane concentrations (< 6 nM) suggesting that the amount of methane flowing back into the Wadden Sea is rather low. A similar observation was found during a tidal inlet study in the German Wadden Sea (Grunwald et al., 2009). Though overall methane concentrations were higher, methane concentrations in North Sea waters flowing into the Wadden Sea were 60 % lower compared to waters flowing out of the Wadden Sea at low tide. Excluding allochthonous methane sources (for example methane influx with freshwater from Lake IJsel), the Dutch Wadden Sea's methane budget must be supported by a total rate of methanogenesis that at least equals the sum of methane efflux to the atmosphere, water column methanotrophy and methane outflow to the North Sea; together these amount to 4.9×10^5 mol d⁻¹. This is comparable to methanogenesis rates in the Eckernförde Bay in the Baltic Sea in the Baltic Sea (Maltby et al., 2018). Note that this accounts for the amount of methane liberated from sediments, while it neglects methane oxidation in sediments (dominantly anaerobic oxidation of methane), which can retain a substantial fraction of methane in sediments (Reeburgh, 2007). Hence, the total rate of methanogenesis in the Wadden Sea is consequently much higher.

Taken all methane export terms/sinks considered together (MOx, efflux and tidal displacement amounting to 4.9×10^5 mol d⁻¹), MOx reduces roughly 2 % of the Wadden Sea's methane budget, while about $1/3^{rd}$ of methane escapes to the atmosphere and the remaining $\sim 2/3^{rd}$ is flushed into the North Sea (where it may be further oxidised and/or released to the atmosphere). The effect of MOx on the methane budget is low when compared to the global ocean, where an estimated >90 % of water column methane is consumed by MOx (Reeburgh, 2007). As the Wadden Sea is very shallow, liberation of methane from sediments to the atmosphere is fast; in other words, MOBs





- have a very limited time to consume methane released from the sediments before it is liberated to the atmosphere
- 477 or flushed with tides to the North Sea. Weber et al. (2019) estimated that coastal systems globally contribute
- between 0.8 and $3.8\,\mathrm{Tg}\,\mathrm{y}^{-1}$ of methane to the atmospheric budget. We found a total annual sea-air flux from the
- Dutch sector of the Wadden Sea (2200 km²) of ~ 0.001 Tg y⁻¹. The Dutch sector of the Wadden Sea alone may
- thus already account for 0.03 % to 0.1 % of the global methane emission from the global coastal ocean.

481 5 Summary and conclusion.

482 Our work revealed substantial variations in methane dynamics when comparing colder and warmer seasons; in 483 warmer seasons, methane concentrations, efflux and MOx were higher compared to colder seasons. Still during 484 colder seasons waters were continuously supersaturated with methane and higher wind speeds in these seasons led 485 to substantial amounts of methane released to the atmosphere. We show that tidal dynamics are a key control for 486 methanotrophic activity and methane distribution. Although changing water column properties and methane 487 concentrations do not provide continuity, the capacity of the microbial methane filter is seemingly stable, with an 488 active MOB community even under unfavourable conditions. Nevertheless, MOx only consumes a minor fraction 489 of the methane inventory of the highly dynamic Wadden Sea, while most is or liberated to the atmosphere and 490 flushed out with tidal currents into the neighbouring North Sea. It appears likely that the contribution of the 491 Wadden Sea to the global atmospheric methane budget will alter in the future due to global warming, and changes 492 in nutrient availability and more frequently occurring storm events. Our results finally highlight the importance of 493 repeated high frequency sampling strategies in dynamic coastal waters to resolve temporal patterns on diel and 494 seasonal scales.

495 Data availability. All data will be archived and made publicly available in the data base DAS (Data Archive
 496 System, www.nioz.nl/en/research/dataportal/das).

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The supplement related to this article is available online.

Author contributions. The study was designed by Tim de Groot, Thomas Röckmann, and Helge Niemann. On board sampling was performed by Tim de Groot, Anne Mol, Katherine Mesdag, Julia Engelmann, Pierre Ramond,
 and Helge Niemann. Further geochemical analysis was conducted by Tim de Groot, Anne Mol, Katherine Mesdag,

and Rachel Ndhlovu. Microbial rates were measured by Tim de Groot and Anne Mol. Statistical analysis was
 carried out by Tim de Groot and Pierre Ramond. Helge Niemann supervised the research project. The manuscript

was prepared by Tim de Groot with input from all authors.

506 Competing interests. The authors disclose that at least one of the (co-)authors holds a position on the editorial507 board of Biogeosciences.

507 board of Biogeoscience

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