1	We would like to thank the editor for the recommendations, which we think helped to improve the
2	quality and clarity of this manuscript. We hope our responses and adaptations are adequate to
3	accept this manuscript for publication in Biogeosciences. Please find our detailed responses below.
4 5 6 7	Associate Editor Comment (8. November 2017)
, 8 9	Dear authors,
10 11 12	I checked your modifications and I am satisfied by your efforts of synthesis. I have some additional recommendations:
13 14 15	- Your manuscript includes too many items (figures and tables). Please select six items that will be presented in the main text and prepare supplementary files for the other items.
15 16 17 18	- Your discussion has too many subtitles. Could you simplify the structuration by, for example, including some titles in the paragraph? To make it visible you can format it in bold or italic.
19 20	Thank you for having submitted to Biogeosciences this nice piece of work.
21 22	Sébastien
23 24 25 26 27 28	Authors Reply: - Even though we are aware that we have many figures, we do think that all of them are necessary, which is why we would like to keep them in the manuscript instead of moving some to the supplement.
29 30 31 32	- We deleted the subheadings (3-level) in the discussion part, as well as the 4-level subheadings in the Material and Methods Part. Instead, we converted them into titles in bold and italic to make the separation between paragraphs visible.
33 34 35	- We added one reference in the paper (introduction part, line 105)
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41	Microbial methanogenesis in the sulfate-reducing zone of sediments in
42	the Eckernförde Bay, SW Baltic Sea
43 44	Johanna Maltby ^{a,b*} , Lea Steinle ^{c,a} , Carolin R. Löscher ^{d,a} , Hermann W. Bange ^a , Martin A. Fischer ^e , Mark Schmidt ^a , Tina Treude ^{f,g*}
45 46	^a GEOMAR Helmholtz Centre for Ocean Research Kiel, Department of Marine Biogeochemistry, 24148 Kiel, Germany
47	^b Present Address: Natural Sciences Department, Saint Joseph's College, Standish, Maine 04084, USA
48	^c Department of Environmental Sciences, University of Basel, 4056 Basel, Switzerland
49	^d Nordic Center for Earth Evolution, University of Southern Denmark, 5230 Odense, Denmark
50	^e Institute of Microbiology, Christian-Albrecht-University Kiel, 24118 Kiel, Germany
51 52	^f Department of Earth, Planetary, and Space Sciences, University of California Los Angeles (UCLA), Los Angeles, California 90095-1567, USA
53 54	g Department of Atmospheric and Oceanic Sciences, University of California Los Angeles (UCLA), Los Angeles, California 90095-1567, USA
56 57	*Correspondence: jmaltby@sjcme.edu, ttreude@g.ucla.edu
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67 Abstract

Benthic microbial methanogenesis is a known source of methane in marine systems. In most 68 69 sediments, the majority of methanogenesis is located below the sulfate-reducing zone, as sulfate 70 reducers outcompete methanogens for the major substrates hydrogen and acetate. Coexistence of 71 methanogenesis and sulfate reduction has been shown before and is possible by usage of non-72 competitive substrates by the methanogens such as methanol or methylated amines. However, the 73 knowledge about magnitude, seasonality and environmental controls on this non-competitive 74 methane production is sparse. In the present study, the presence of methanogenesis within the 75 sulfate reduction zone (SRZ methanogenesis), was investigated in sediments (0-30 centimeters below 76 seafloor, cmbsf) of the seasonally hypoxic Eckernförde Bay, southwestern Baltic Sea. Water column 77 parameters such as oxygen, temperature, and salinity together with porewater geochemistry and 78 benthic methanogenesis rates were determined in the sampling area "Boknis Eck" quarterly from 79 March 2013 to September 2014, to investigate the effect of seasonal environmental changes on the 80 rate and distribution of SRZ methanogenesis, to estimate its potential contribution to benthic 81 methane emissions, and to identify potential methanogenic groups responsible for SRZ 82 methanogenesis. The metabolic pathway of methanogenesis in the presence or absence of sulfate reducers and after the addition of a non-competitive substrate was studied in four experimental 83 setups: 1) unaltered sediment batch incubations (net methanogenesis), 2) ¹⁴C-bicarbonate labeling 84 85 experiments (hydrogenotrophic methanogenesis), 3) manipulated experiments with addition of 86 either molybdate (sulfate reducer inhibitor), 2-bromoethane-sulfonate (methanogen inhibitor), or methanol (non-competitive substrate, potential methanogenesis), 4) addition of ¹³C-labeled 87 88 methanol (potential methylotrophic methanogenesis). After incubation with methanol, molecular 89 analyses were conducted to identify key functional methanogenic groups during methylotrophic 90 methanogenesis. To also compare magnitudes of SRZ methanogenesis with methanogenesis below 91 the sulfate reduction zone (> 30 cmbsf), hydrogenotrophic methanogenesis was determined by ¹⁴Cbicarbonate radiotracer incubation in samples collected in September 2013. 92 93 SRZ methanogenesis changed seasonally in the upper 30 cmbsf with rates increasing from March (0.2 nmol cm⁻³ d⁻¹) to November (1.3 nmol cm⁻³ d⁻¹) 2013 and March (0.2 nmol cm⁻³ d⁻¹) to September (0.4 94 95 nmol cm⁻³ d⁻¹) 2014, respectively. Its magnitude and distribution appeared to be controlled by 96 organic matter availability, C/N, temperature, and oxygen in the water column, revealing higher rates 97 in warm, stratified, hypoxic seasons (September/November) compared to colder, oxygenated 98 seasons (March/June) of each year. The majority of SRZ methanogenesis was likely driven by the usage of non-competitive substrates (e.g., methanol and methylated compounds), to avoid 99 100 competition with sulfate reducers, as it was indicated by the 1000-3000-fold increase in potential 101 methanogenesis activity observed after methanol addition. Accordingly, competitive

- 102 hydrogenotrophic methanogenesis increased in the sediment only below the depth of sulfate
- 103 penetration (> 30 cmbsf). Members of the family Methanosarcinaceae, which are known for
- 104 methylotrophic methanogenesis, were detected by PCR using *Methanosarcinaceae*-specific primers
- and are likely to be responsible for the observed SRZ methanogenesis.
- 106 The present study indicates that SRZ methanogenesis is an important component of the benthic
- 107 methane budget and carbon cycling in Eckernförde Bay. Although its contribution to methane
- 108 emissions from the sediment into the water column are probably minor, SRZ methanogenesis could
- 109 directly feed into methane oxidation above the sulfate-methane transition zone.

110 1. Introduction

111 After water vapor and carbon dioxide, methane is the most abundant greenhouse gas in the 112 atmosphere (e.g. Hartmann et al., 2013; Denman et al., 2007). Its atmospheric concentration 113 increased more than 150 % since preindustrial times, mainly through increased human activities such 114 as fossil fuel usage and livestock breeding (Hartmann et al., 2013; Wuebbles & Hayhoe, 2002; 115 Denman et al., 2007). Determining the natural and anthropogenic sources of methane is one of the 116 major goals for oceanic, terrestrial and atmospheric scientists to be able to predict further impacts 117 on the world's climate. The ocean is considered to be a modest natural source for atmospheric 118 methane (Wuebbles & Hayhoe, 2002; Reeburgh, 2007; EPA, 2010). However, research is still sparse 119 on the origin of the observed oceanic methane, which automatically leads to uncertainties in current 120 ocean flux estimations (Bange et al., 1994; Naqvi et al., 2010; Bakker et al., 2014). Within the marine environment, the coastal areas (including estuaries and shelf regions) are 121 122 considered the major source for atmospheric methane, contributing up to 75 % to the global ocean 123 methane production (Bange et al., 1994). The major part of the coastal methane is produced during 124 microbial methanogenesis in the sediment, with probably only a minor part originating from 125 methane production within the water column (Bakker et al., 2014). However, the knowledge on 126 magnitude, seasonality and environmental controls of benthic methanogenesis is still limited. 127 In marine sediments, methanogenesis activity is mostly restricted to the sediment layers below 128 sulfate reduction, due to the successful competition of sulfate reducers with methanogens for the 129 mutual substrates acetate and hydrogen (H₂) (Oremland & Polcin, 1982; Crill & Martens, 1986; 130 Jørgensen, 2006). Methanogens produce methane mainly from using acetate (acetoclastic methanogenesis) or H₂ and carbon dioxide (CO₂) (hydrogenotrophic methanogenesis). Competition 131 132 with sulfate reducers can be relieved through usage of non-competitive substrates (e.g. methanol or 133 methylated compounds, methylotrophic methanogenesis) (Cicerone & Oremland, 1988; Oremland & 134 Polcin, 1982). Coexistence of sulfate reduction and methanogenesis has been detected in a few 135 studies from organic-rich sediments, e.g., salt-marsh sediments (Oremland et al., 1982; Buckley et al.,

2008), coastal sediments (Holmer & Kristensen, 1994; Jørgensen & Parkes, 2010) or sediments in 136 137 upwelling regions (Pimenov et al., 1993; Ferdelman et al., 1997; Maltby et al., 2016), indicating the 138 importance of these environments for methanogenesis within the sulfate reduction zone (SRZ 139 methanogenesis). So far, however, environmental controls of SRZ methanogenesis remain elusive. The coastal inlet Eckernförde Bay (southwestern Baltic Sea) is an excellent model environment to 140 study seasonal and environmental controls of benthic SRZ methanogenesis. Here, the muddy 141 142 sediments are characterized by high organic loading and high sedimentation rates (Whiticar, 2002), 143 which lead to anoxic conditions within the uppermost 0.1-0.2 centimeter below seafloor (cmbsf) 144 (Preisler et al., 2007). Seasonally hypoxic (dissolved oxygen < 63 µM) and anoxic (dissolved oxygen = 145 0 µM) events in the bottom water of Eckernförde Bay (Lennartz et al., 2014; Steinle et al., 2017) 146 provide ideal conditions for anaerobic processes at the sediment surface. 147 Sulfate reduction is the dominant pathway of organic carbon degradation in Eckernförde Bay 148 sediments in the upper 30 cmbsf, followed by methanogenesis in deeper sediment layers where 149 sulfate is depleted (>> 30 cmbsf) (Whiticar 2002; Treude et al. 2005; Martens et al. 1998) (Fig. 1). This 150 methanogenesis below the sulfate-methane transition zone (SMTZ) can be intense and often leads to methane oversaturation in the porewater below 50 cm sediment depth, resulting in gas bubble 151 formation (Abegg & Anderson, 1997; Whiticar, 2002; Thießen et al., 2006). Thus, methane is 152 153 transported from the methanogenic zone (> 30 cmbsf) to the surface sediment by both molecular 154 diffusion and advection via rising gas bubbles (Wever et al., 1998; Treude et al., 2005a). Although 155 upward diffusing methane is mostly retained by anaerobic oxidation of methane (AOM) (Treude et 156 al. 2005), a major part is reaching the sediment-water interface through gas bubble transport 157 (Treude et al. 2005; Jackson et al. 1998), resulting in a supersaturation of the water column with respect to atmospheric methane concentrations (Bange et al., 2010). The Time Series Station "Boknis 158 Eck" in the Eckernförde Bay is a known site of methane emissions into the atmosphere throughout 159 160 the year due to this supersaturation of the water column (Bange et al., 2010). 161 The source for benthic and water column methane was seen in methanogenesis below the SMTZ (>> 162 30 cmbsf) (Whiticar, 2002), however, coexistence of sulfate reduction and methanogenesis has been 163 postulated (Whiticar, 2002; Treude et al., 2005a). Still, the magnitude and environmental controls of 164 SRZ methanogenesis is poorly understood, even though it may make a measurable contribution to 165 benthic methane emissions given its short diffusion distance to the sediment-water interface (Knittel 166 & Boetius, 2009). Production of methane within the sulfate reduction zone of Eckernförde Bay 167 sediments could further explain peaks of methane oxidation observed in top sediment layers, which was previously attributed to methane transported to the sediment surface via rising gas bubbles 168 169 (Treude et al., 2005a).

170 In the present study, we investigated sediments from within (< 30 cmbsf, on a seasonal basis) and 171 below the sulfate reduction zone (>> 30 cmbsf, on one occasion), and the water column (on a 172 seasonal basis) at the Time Series Station "Boknis Eck" in Eckernförde Bay, to validate the existence 173 of SRZ methanogenesis and its potential contribution to benthic methane emissions. Water column parameters like oxygen, temperature, and salinity together with porewater geochemistry and 174 175 benthic methanogenesis were measured over a course of 2 years. In addition to seasonal rate 176 measurements, inhibition and stimulation experiments, stable isotope probing, and molecular 177 analysis were carried out to find out if SRZ methanogenesis 1) is controlled by environmental 178 parameters, 2) shows seasonal variability, 3) is based on non-competitive substrates with a special 179 focus on methylotrophic methanogens.

180 2. Material and Methods

181 2.1 Study site

182 Samples were taken at the Time Series Station "Boknis Eck" (BE, 54°31.15 N, 10°02.18 E; 183 www.bokniseck.de) located at the entrance of Eckernförde Bay in the southwestern Baltic Sea with a 184 water depth of about 28 m (map of sampling site can be found in e.g. Hansen et al., (1999)). From 185 mid of March until mid of September the water column is strongly stratified due to the inflow of 186 saltier North Sea water and a warmer and fresher surface water (Bange et al., 2011). Organic matter degradation in the deep layers causes pronounced hypoxia (March-Sept) or even anoxia 187 (August/September) (Smetacek, 1985; Smetacek et al., 1984). The source of organic material is 188 189 phytoplankton blooms that occur regularly in spring (February-March) and fall (September-190 November) and are followed by pronounced sedimentation of organic matter (Bange et al., 2011). To 191 a lesser extent, phytoplankton blooms and sedimentation are also observed during the summer 192 months (July/August) (Smetacek et al., 1984). Sediments at BE are generally classified as soft, fine-193 grained muds (< 40 μm) with a carbon content of 3 to 5 wt% (Balzer et al., 1986). The bulk of organic matter in Eckernförde Bay sediments originates from marine plankton and macroalgal sources (Orsi 194 195 et al., 1996), and its degradation leads to production of free methane gas (Wever & Fiedler, 1995; 196 Abegg & Anderson, 1997; Wever et al., 1998). The oxygen penetration depth is limited to the upper 197 few millimeters when bottom waters are oxic (Preisler et al., 2007). Reducing conditions within the 198 sulfate reduction zone lead to a dark grey/black sediments color with a strong hydrogen sulfur odor 199 in the upper meter of the sediment and dark olive-green color the deeper sediment layers (> 1 m) 200 (Abegg & Anderson, 1997).

201 2.2 Water column and sediment sampling

Sampling was done on a seasonal basis during the years of 2013 and 2014. One-Day field trips with 202 203 either F.S. Alkor (cruise no. AL410), F.K. Littorina or F.B. Polarfuchs were conducted in March, June, 204 and September of each year. In 2013, additional sampling was conducted in November. At each sampling month, water profiles of temperature, salinity, and oxygen concentration (optical sensor, 205 206 RINKO III, detection limit= 2 µM) were measured with a CTD (Hydro-Bios). In addition, water samples 207 for methane concentration measurements were taken at 25 m water depth with a 6-Niskin bottle (4 208 Liter each) rosette attached to the CTD (Table 1). Complementary samples for water column 209 chlorophyll were taken at 25 m water depth with the CTD-rosette within the same months during 210 standardized monthly sampling cruises to Boknis Eck organized by GEOMAR. 211 Sediment cores were taken with a miniature multicorer (MUC, K.U.M. Kiel), holding 4 core liners 212 (length= 60 cm, diameter= 10 cm) at once. The cores had an average length of ~ 30 cm and were 213 stored at 10°C in a cold room (GEOMAR) until further processing (normally within 1-3 days after 214 sampling).

- 215 In September 2013, a gravity core was taken in addition to the MUC cores. The gravity core was
- 216 equipped with an inner plastic bag (polyethylene; diameter: 13 cm). After core recovery (330 cm
- total length), the polyethylene bag was cut open at 12 different sampling depths resulting in intervals
- of 30 cm and sampled directly on board for sediment porewater geochemistry (see Sect. 2.4),
- 219 sediment methane (see Sect. 2.5), sediment solid phase geochemistry (see Sect. 2.6), and microbial
- rate measurements for hydrogenotrophic methanogenesis as described in section 2.8.

221 2.3 Water column parameters

- At each sampling month, water samples for methane concentration measurements were taken at 25
- 223 m water depth in triplicates. Therefore, three 25 ml glass vials were filled bubble free directly after
- 224 CTD-rosette recovery and closed with butyl rubber stoppers. Biological activity in samples was
- 225 stopped by adding saturated mercury chloride solution, followed by storage at room temperature
- 226 until further treatment.
- 227 Concentrations of dissolved methane (CH₄) were determined by headspace gas chromatography as
- 228 described in Bange et al. (2010). Calibration for CH₄ was done by a two-point calibration with known
- 229 methane concentrations before the measurement of headspace gas samples, resulting in an error of
 230 < 5 %.</p>
- 231 Water samples for chlorophyll concentration were taken by transferring the complete water volume
- 232 (from 25 m water depth) from one water sampler into a 4.5 L Nalgene bottle, from which then
- 233 approximately 0.7-1 L (depending on the plankton content) were filtrated back in the GEOMAR
- 234 laboratory using GF/F filter (Whatman, 25 mm diameter, 8 μM pores size). Dissolved chlorophyll a

concentrations were determined using the fluorometric method by Welschmeyer (1994) with an
 error < 10 %.

237 2.4 Sediment porewater geochemistry

- 238 Porewater was extracted from sediment within 24 hours after core retrieval using nitrogen (N₂) pre-
- 239 flushed rhizons (0.2 μm, Rhizosphere Research Products, Seeberg-Elverfeldt et al., 2005). In MUC
- 240 cores, rhizons were inserted into the sediment in 2 cm intervals through pre-drilled holes in the core
- liner. In the gravity core, rhizons were inserted into the sediment in 30 cm intervals directly afterretrieval.
- 243 Extracted porewater from MUC and gravity cores was immediately analyzed for sulfide using
- standardized photometric methods (Grasshoff et al., 1999).
- 245 Sulfate concentrations were determined using ion chromatography (Methrom 761). Analytical
- 246 precision was < 1 % based on repeated analysis of IAPSO seawater standards (dilution series) with an
- absolute detection limit of 1 μ M corresponding to a detection limit of 30 μ M for the undiluted sample.
- 249 For analysis of dissolved inorganic carbon (DIC), 1.8 ml of porewater was transferred into a 2 ml glass
- 250 vial, fixed with 10 μ l saturated HgCL₂ solution and crimp sealed. DIC concentration was determined
- as CO₂ with a multi N/C 2100 analyzer (Analytik Jena) following the manufacturer's instructions.
- 252 Therefore, the sample was acidified with phosphoric acid and the outgassing CO₂ was measured. The
- 253 detection limit was 20 μM with a precision of 2-3 %.

254 2.5 Sediment methane concentrations

- 255 In March 2013, June 2013 and March 2014, one MUC core was sliced in 1 cm intervals until 6 cmbsf,
- followed by 2 cm intervals until the end of the core. At the other sampling months, the MUC core
- was sliced in 1 cm intervals until 6 cmbsf, followed by 2 cm intervals until 10 cmbsf and 5 cm intervalsuntil the end of the core.
- 259 Per sediment depth (in MUC and gravity cores), 2 cm⁻³ of sediment were transferred into a 10 ml-
- 260 glass vial containing 5 ml NaOH (2.5 %) for determination of sediment methane concentration per
- 261 volume of sediment. The vial was quickly closed with a butyl septum, crimp-sealed and shaken
- 262 thoroughly. The vials were stored upside down at room temperature until measurement via gas
- 263 chromatography. Therefore, 100 µl of headspace was removed from the gas vials and injected into a
- 264 Shimadzu gas chromatograph (GC-2014) equipped with a packed Haysep-D column and a flame
- 265 ionization detector. The column temperature was 80°C and the helium flow was set to 12 ml min⁻¹.
- 266 CH₄ concentrations were calibrated against CH₄ standards (Scotty gases). The detection limit was 0.1
- 267 ppm with a precision of 2 %.

268 2.6 Sediment solid phase geochemistry

- 269 Following the sampling for CH₄, the same cores described under section 2.5 were used for the
- 270 determination of the sediment solid phase geochemistry, i.e. porosity, particulate organic carbon
- 271 (POC) and particulate organic nitrogen (PON).
- 272 Sediment porosity of each sampled sediment section was determined by the weight difference of 5
- 273 cm⁻³ wet sediment after freeze-drying for 24 hours. Dried sediment samples were then used for
- analysis of particulate organic carbon (POC) and particulate organic nitrogen (PON) with a Carlo-Erba
- element analyzer (NA 1500). The detection limit for C and N analysis was < 0.1 dry weight percent (%)
 with a precision of < 2 %.

277 2.7 Sediment methanogenesis

278 2.7.1 Methanogenesis in MUC cores

- 279 At each sampling month, three MUC cores were sliced in 1 cm intervals until 6 cmbsf, in 2 cm
- intervals until 10 cmbsf, and in 5 cm intervals until the bottom of the core. Every sediment layer was
- 281 transferred to a separate beaker and quickly homogenized before sub-sampling. The exposure time
- 282 with air, i.e. oxygen, was kept to a minimum. Sediment layers were then sampled for determination
- 283 of net methanogenesis (defined as the sum of total methane production and consumption, including
- all available methanogenic substrates in the sediment), hydrogenotrophic methanogenesis
- 285 (methanogenesis based on the substrates CO₂/H₂), and potential methanogenesis (methanogenesis
- at ideal conditions, i.e. no lack of nutrients) as described in the following sections.

287 2.7.1.1 Net methanogenesis

- Net methanogenesis was determined with sediment slurry experiments by measuring the headspace
 methane concentration over time. Per sediment layer, triplicates of 5 cm⁻³ of sediment were
- 290 transferred into N₂-flushed sterile glass vials (30 ml) and mixed with 5 ml filtered bottom water. The
- slurry was repeatedly flushed with N₂ to remove residual methane and to ensure complete anoxia.
- 292 Slurries were incubated in the dark at in-situ temperature, which varied at each sampling date (Table
- 1). Headspace samples (0.1 ml) were taken out every 3-4 days over a time period of 4 weeks and
- analyzed on a Shimadzu GC-2104 gas chromatograph (see Sect. 2.5). Net methanogenesis rates were
- 295 determined by the linear increase of the methane concentration over time (minimum of 6 time
- 296 points, see also Fig. S1).

297 2.7.1.2 Hydrogenotrophic methanogenesis 298 To determine if hydrogenotrophic methanogenesis, i.e., methanogenesis based on the competitive 299 substrates H₂, is present in the sulfate-reducing zone, radioactive sodium bicarbonate (NaH¹⁴CO₃)

300 was added to the sediment.

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- 301 Per sediment layer, sediment was sampled in triplicates with glass tubes (5 mL) which were closed
- 302 with butyl rubber stoppers on both ends according to (Treude et al. 2005). Through the stopper,
- 303 NaH¹⁴CO₃ (dissolved in water, injection volume 6 μ l, activity 222 kBq, specific activity = 1.85-2.22
- 304 GBq/mmol) was injected into each sample and incubated for three days in the dark at in-situ
- 305 temperature (Table 1). To stop bacterial activity, sediment was transferred into 50 ml glass-vials filled
- 306 with 20 ml sodium hydroxide (2.5 % w/w), closed quickly with rubber stoppers and shaken
- thoroughly. Five controls were produced from various sediment depths by injecting the radiotracerdirectly into the NaOH with sediment.
- 309 The production of ¹⁴C-methane was determined with the slightly modified method by Treude et al.,
- (2005) used for the determination of anaerobic oxidation of methane. The method was identical,
- 311 except no unlabeled methane was determined by gas chromatography. Instead, DIC values were
- 312 used to calculate hydrogenotrophic methane production.

313 **2.7.1.3** Potential methanogenesis in manipulated experiments

- To examine the interaction between sulfate reduction and methanogenesis, inhibition and
- 315 stimulation experiments were carried out. Therefore, every other sediment layer was sampled
- resulting in the following examined six sediment layers: 0-1 cm, 2-3 cm, 4-5 cm, 6-8 cm, 10-15 cm
- and 20-25 cm. From each layer, sediment slurries were prepared by mixing 5 ml sediment in a 1:1
- ratio with adapted artificial seawater medium (salinity 24, Widdel & Bak, 1992) in N₂-flushed, sterile
- 319 glass vials before further manipulations.
- 320 In total, four different treatments, each in triplicates, were prepared per depth: 1) with sulfate
- 321 addition (17 mM), 2) with sulfate (17 mM) and molybdate (22 mM) addition, 3) with sulfate (17 mM)
- 322 and 2-bromoethane-sulfonate (BES, 60 mM) addition, and 4) with sulfate (17 mM) and methanol (10
- 323 mM) addition. From here on, the following names are used to describe the different treatments,
- 324 respectively: 1) control treatment, 2) molybdate treatment, 3) BES treatment, and 4) methanol
- 325 treatment. Control treatments feature the natural sulfate concentrations occurring in sediments of
- 326 the sulfate reduction zone at the sampling site. Molybdate was used as an enzymatic inhibitor for
- 327 sulfate reduction (Oremland & Capone, 1988) and BES was used as an inhibitor for methanogenic
- 328 archaea (Hoehler et al., 1994). Methanol is a known non-competitive substrate, which is used by
- 329 methanogens but not by sulfate reducers (Oremland & Polcin, 1982), thus it is suitable to examine
- 330 non-competitive methanogenesis. Treatments were incubated similar to net methanogenesis
- 331 (2.7.1.1) by incubating sediment slurries at the respective in-situ temperature (Table 1) in the dark
- for a time period of 4 weeks. Headspace samples (0.1 ml) were taken every 3-5 days over a time
- 333 period of 4 weeks and potential methanogenesis rates were determined by the linear increase of
- 334 methane concentration over time (minimum of 6 time points).

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336	2.7.1.4 Potential methylotrophic methanogenesis from methanol using stable isotope probing	
337	One additional experiment was conducted with sediments from September 2014 by adding 13 C-	
338	labelled methanol to investigate the production of ¹³ C-labelled methane. Three cores were stored at	
339	1°C after the September 2014 cruise until further processing \sim 3.5 months later. The low storage	
340	temperature together with the expected oxygen depletion in the enclosed supernatant water after	
341	retrieval of the cores likely led to slowed anaerobic microbial activity during storage time and	
342	preserved the sediments for potential methanogenesis measurements.	
343	Sediment cores were sliced in 2 cm intervals and the upper 0-2 cmbsf sediment layer of all three	
344	cores was combined in a beaker and homogenized. Then, sediment slurries were prepared by mixing	
345	5 cm $^{-3}$ of sediment with 5 ml of artificial seawater medium in N2-flushed, sterile glass vials (30 ml).	
346	After this, methanol was added to the slurry with a final concentration of 10 mM (see 2.7.1.3).	
347	Methanol was enriched with 13 C-labelled methanol in a ratio of 1:1000 between 13 C-labelled (99.9 %	
348	13 C) and non-labelled methanol mostly consisting of 12 C (manufacturer: Roth). In total, 54 vials were	
349	prepared for nine different sampling time points during a total incubation time of 37 days. All vials	
350	were incubated at 13°C (in situ temperature in September 2014) in the dark. At each sampling point,	
351	six vials were stopped: one set of triplicates were used for headspace methane and carbon dioxide	
352	determination and a second set of triplicates were used for porewater analysis.	
353	Headspace methane and carbon dioxide concentrations (volume 100 $\mu l)$ were determined on a	
354	Shimadzu gas chromatograph (GC-2014) equipped with a packed Haysep-D column a flame ionization	
355	detector and a methanizer. The methanizer (reduced nickel) reduces carbon dioxide with hydrogen	
356	to methane at a temperature of 400°C. The column temperature was 80°C and the helium flow was	
357	set to 12 ml min $^{-1}$. Methane concentrations (including reduced CO ₂) were calibrated against methane	
358	standards (Scotty gases). The detection limit was 0.1 ppm with a precision of 2 %.	
359	Analyses of ¹³ C/ ¹² C-ratios of methane and carbon dioxide were conducted after headspace	
360	concentration measurements by using a continuous flow combustion gas chromatograph (Trace	
361	Ultra, Thermo Scientific), which was coupled to an isotope ratio mass spectrometer (MAT253,	
362	Thermo Scientific). The isotope ratios of methane and carbon dioxide given in the common delta-	
363	notation (S $^{\rm 13}{\rm C}$ in permill) are reported relative to Vienna Pee Dee Belemnite (VPDB) standard.	
364	Isotope precision was +/- 0.5 ‰, when measuring near the detection limit of 10 ppm.	
365	For porewater analysis of methanol concentration and isotope composition, each sediment slurry of	
366	the triplicates was transferred into argon-flushed 15 ml centrifuge tubes and centrifuged for 6	
367	minutes at 4500 rpm. Then 1 ml filtered (0.2 μm) porewater was transferred into N2-flushed 2 ml	
368	glass vials for methanol analysis, crimp sealed and immediately frozen at -20 °C. Methanol	

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- 369 concentrations and isotope composition were determined via high performance liquid
- 370 chromatography-ion ratio mass spectrometry (HPLC-IRMS, Thermo Fisher Scientific) at the MPI
- 371 Marburg. The detection limit was 50 μ M with a precision of 0.3‰.

372 2.7.2 Methanogenesis in the gravity core

- 373 Ex situ hydrogenotrophic methanogesis was determined in a gravity core taken in September 2013.
- 374 The pathway is thought to be the main methanogenic pathway in the sediment below the SMTZ in
- 375 Eckernförde Bay (Whiticar, 2002). Hydrogenotrophic methanogenesis was determined using
- 376 radioactive sodium bicarbonate (NaH¹⁴CO₃). At every sampled sediment depth (12 depths in 30 cm
- 377 intervals), triplicate glass tubes (5 mL) were inserted directly into the sediment. Tubes were filled
- 378 bubble-free with sediment and closed with butyl rubber stoppers on both ends according to (Treude
- et al. 2005). Methods following sampling were identical as described in 2.7.1.2.

380 2.8 Molecular analysis

- 381 During the non-labeled methanol treatment of the 0-1 cmbsf horizon from the September 2014 382 sampling (see 2.7.1.3), additional samples were prepared to detect and quantify the presence of methanogens in the sediment. Therefore, additional 15 vials were prepared with addition of 383 methanol as described in 2.7.1.3 for five different time points (day 1 (= t_0), day 8, day 16, day 22, and 384 385 day 36) and stopped at each time point by transferring sediment from the triplicate slurries into 386 whirl-packs (Nasco), which then were immediately frozen at -20°C. DNA was extracted from ~500 mg of sediment using the FastDNA® SPIN Kit for Soil (Biomedical). Quantitative real-time polymerase 387 chain reaction (qPCR) technique using TaqMan probes and TaqMan chemistry (Life Technologies) was 388 used for the detection of methanogens on a ViiA7 qPCR machine (Life Technologies). Primer and 389 390 Probe sets as originally published by Yu et al. (2005) were applied to quantify the orders Methanobacteriales, Methanosarcinales and Methanomicrobiales along with the two families 391 392 Methanosarcinaceae and Methanosaetaceae within the order Methanosarcinales. In addition, a 393 universal primer set for detection of the domain Archaea was used (Yu et al. 2005). Absolut quantification of the 16S rDNA from the groups mentioned above was performed with 394 395 standard dilution series. The standard concentration reached from 10⁸ to 10¹ copies per µL. Quantification of the standards and samples was performed in duplicates. Reaction was performed in 396 397 a final volume of 12.5 μ L containing 0.5 μ L of each Primer (10pmol μ L⁻¹, MWG), 0.25 μ L of the 398 respective probe (10 pmol μ L⁻¹, Life Technologies), 4 μ L H₂O (Roth), 6.25 μ L TaqMan Universal Master Mix II (Life Technologies) and 1 µL of sample or standard. Cycling conditions started with initial 399 400 denaturation and activation step for 10 min at 95°C, followed by 45 cycles of 95 °C for 15 sec, 56°C 401 for 30 sec and 60°C for 60 sec. Non-template controls were run in duplicates with water instead of
- 402 DNA for all primer and probe sets, and remained without any detectable signal after 45 cycles.

403 2.9 Statistical Analysis

- 404 To determine possible environmental control parameters of SRZ methanogenesis, a Principle 405 Component Analysis (PCA) was applied according to the approach described in Gier et al. (2016). 406 Prior to PCA, the dataset was transformed into ranks to assure the same data dimension. In total, two PCAs were conducted. The first PCA was used to test the relation of parameters in the 407 408 surface sediment (integrated methanogenesis (0-5 cm, mmol m⁻² d⁻¹), POC content (average value 409 from 0-5 cmbsf, wt %), C/N (average value from 0-5 cmbsf, molar) and the bottom water (25 m water 410 depth) (oxygen (µM), temperature (°C), salinity (PSU), chlorophyll (µg L-1), methane (nM)). The 411 second PCA was applied on depth profiles of sediment SRZ methanogenesis (nmol cm⁻³ d⁻¹), sediment 412 depth (cm), sediment POC content (wt%), sediment C/N ratio (molar), and sampling month (one 413 value per depth profile at a specific month, the later in the year the higher the value). 414 For each PCA, biplots were produced to view data from different angles and to graphically determine 415 a potential positive, negative or zero correlation between methanogenesis rates and the tested 416 variables.
- 417 3. Results

418 3.1 Water column parameters

- 419 From March 2013 to September 2014, the water column had a pronounced temporal and spatial
- 420 variability of temperature, salinity, and oxygen (Fig. 2 and 3). In 2013, temperature of the upper
- 421 water column increased from March (1°C) to September (16°C), but decreased again in November
- 422 (11°C). The temperature of the lower water column increased from March 2013 (2°C) to November
- 423 2013 (12°C). In 2014, lowest temperatures of the upper and lower water column were reached in
- 424 March (4°C). Warmer temperatures of the upper water column were observed in June and
- 425 September (around 17°C), while the lower water column peaked in September (13°C).
- 426 Salinity increased over time during 2013, showing the highest salinity of the upper and lower water
- 427 column in November (18 and 23 PSU, respectively). In 2014, salinity of the upper water column was
- 428 highest in March and September (both 17 PSU), and lowest in June (13 PSU). The salinity of the lower
- 429 water column increased from March 2014 (21 PSU) to September 2014 (25 PSU).
- In both years, June and September showed the most pronounced vertical gradient of temperatureand salinity, featuring a pycnocline at around ~14 m water depth.
- 432 Summer stratification was also seen in the O₂ profiles, which showed O₂ depleted conditions (O₂ <
- 433 150 μM) in the lower water column from June to September in both years, reaching concentrations
- 434 below 1- 2 μ M (detection limit of CTD sensor) in September of both years (Fig. 2 and 3). The water
- 435 column was completely ventilated, i.e. homogenized, in March of both years with O₂ concentrations
- 436 of 300-400 μ M down to the sea floor at about 28 m.

437

438 3.2 Sediment geochemistry in MUC cores 439 Sediment porewater and solid phase geochemistry results for the years 2013 and 2014 are shown in 440 Fig. 2 and 3, respectively. 441 Sulfate concentrations at the sediment surface ranged between 15-20 mM. Concentration decreased with depth at all sampling months but was never fully depleted until the bottom of the core (18-29 442 443 cmbsf, between 2 and 7 mM sulfate). November 2013 showed the strongest decrease from ~20 mM 444 at the top to ~2 mM at the bottom of the core (27 cmbsf). 445 Opposite to sulfate, methane concentration increased with sediment depth in all sampling months 446 (Fig. 2 and 3). Over the course of a year (i.e. March to November in 2013, and March to September in 447 2014), maximum methane concentration increased, reaching the highest concentration in November 448 2013 (~1 mM at 26 cmbsf) and September 2014 (0.2 mM at 23 cmbsf), respectively. Simultaneously, 449 methane profiles became steeper, revealing higher methane concentrations at shallower sediment 450 depth late in the year. Magnitudes of methane concentrations were similar in the respective months 451 of 2013 and 2014. 452 In all sampling months, sulfide concentration increased with sediment depth (Fig. 2 and 3). Similar to 453 methane, sulfide profiles revealed higher sulfide concentrations at shallower sediment depth 454 together with higher peak concentrations over the course the sampled months in each sampling year. Accordingly, November 2013 (10.5 mM at 15 cmbsf) and September 2014 (2.8 mM at 15 455 456 cmbsf) revealed the highest sulfide concentrations, respectively. September 2014 was the only 457 sampling month showing a pronounced decrease in sulfide concentration from 15 cmbsf to 21 cmbsf 458 of over 50 %. 459 DIC concentrations increased with increasing sediment depth at all sampling months. Concomitant with highest sulfide concentrations, highest DIC concentration was detected in November 2013 (26 460 mM at 27 cmbsf). At the surface, DIC concentrations ranged between 2-3 mM at all sampling 461 462 months. In June of both years, DIC concentrations were lowest at the deepest sampled depth compared to the other sampling months (16 mM in 2013, 13 mM in 2014). 463 464 At all sampling months, POC profiles scattered around 5 \pm 0.9 wt % with depth. Only in November 2013, June 2014 and September 2014, POC content exceeded 5 wt % in the upper 0-1 cmbsf (5.9, 5.2 465 466 and 5.3 wt %, respectively) with the highest POC content in November 2013. Also in November 2013, 467 surface C/N ratio (0-1 cmbsf) of the particulate organic matter was lowest of all sampling months

- 468 (8.6). In general, C/N ratio increased with depth in both years with values around 9 at the surface and
- values around 10-11 at the deepest sampled sediment depths.

470 **3.3 Sediment geochemistry in gravity cores**

471 Results from sediment porewater and solid phase geochemistry in the gravity core from September 472 2013 are shown in Fig. 4. Please note that the sediment depth of the gravity core was corrected by 473 comparing the sulfate concentrations at 0 cmbsf in the gravity core with the corresponding sulfate concentration and depth in the MUC core from September 2013 (Fig. 2). The soft surface sediment is 474 475 often lost during the gravity coring procedure. Through this correction, the topmost layer of the 476 gravity core was set at a depth of 14 cmbsf. 477 Porewater sulfate concentration in the gravity core decreased with depth (i.e. below 0.1 mM at 107 478 cmbsf) and stayed below 0.1 mM until 324 cmbsf. Sulfate increased slightly (1.9 mM) at the bottom 479 of the core (345 cmbsf). In concert with sulfate, also methane, sulfide, DIC, POC and C/N profiles 480 showed distinct alteration in the profile at 345 cmbsf (see below, Fig. 4). As fluid seepage has not 481 been observed at the Boknis Eck station (Schlüter et al., 2000), these alterations could either indicate 482 a change in sediment properties or result from a sampling artifact from the penetration of seawater 483 through the core catcher into the deepest sediment layer. The latter process is, however, not 484 expected to considerably affect sediment solid phase properties (POC and C/N), and we therefore 485 dismissed this hypothesis. Methane concentration increased steeply with depth reaching a maximum of 4.8 mM at 76 cmbsf. 486 487 Concentration stayed around 4.7 mM until 262 cmbsf, followed by a slight decrease until 324 cmbsf 488 (2.8 mM). From 324 cmbsf to 345 cmbsf methane increased again (3.4 mM). Both sulfide and DIC concentrations increased with depth, showing a maximum at 45 cmbsf (~ 5mM) 489 490 and 345 cmbsf (~ 1mM), respectively. While sulfide decreased after 45 cmbsf to a minimum of ~ 300

491 $~~\mu\text{M}$ at 324 cmbsf, it slightly increased again to ~1 mM at 345 cmbsf. In accordance, DIC

492 concentrations showed a distinct decrease between 324 cmbsf to 345 cmbsf (from 45 mM to 39493 mM).

494 While POC contents varied around 5 wt % throughout the core, C/N ratio slightly increased with

495 depth, revealing the lowest ratio at the surface (~3) and the highest ratio at the bottom of the core

496 (~13). However, both POC and C/N showed a distinct increase from 324 cmbsf to 345 cmbsf.

497

498 3.4 Methanogenesis activity in MUC cores

499 3.4.1 Net methanogenesis

- 500 Net methanogenesis activity (calculated by the linear increase of methane over time, see Fig. S1) was
- 501 detected throughout the cores at all sampling months (Fig. 2 and 3). Activity measured in MUC cores
- 502 increased over the course of the year in 2013 and 2014 (that is: March to November in 2013 and
- 503 March to September in 2014) with lower rates mostly < 0.1 nmol cm⁻³ d⁻¹ in March and higher rates >

- 504 0.2 nmol cm⁻³ d⁻¹ in November 2013 and September 2014, respectively. In general, November 2013
- 505 revealed highest net methanogenesis rates (1.3 nmol cm⁻³ d⁻¹ at 1-2 cmbsf). Peak rates were
- 506 detected at the sediment surface (0-1 cmbsf) at all sampling months except for September 2013
- 507 where the maximum rates were situated between 10-15 cmbsf. In addition to the surface peaks, net
- 508 methanogenesis showed subsurface (= below 1 cmbsf until 30 cmbsf) maxima at all sampling
- 509 months, but with alternating depths (between 10 and 25 cmbsf).
- 510 Comparison of integrated net methanogenesis rates (0-25 cmbsf) revealed highest rates in
- 511 September and November 2013 (0.09 mmol m⁻² d⁻¹ and 0.08 mmol m⁻² d⁻¹, respectively) and lowest
- 512 rates in March 2014 (0.01 mmol m⁻² d⁻¹) (Fig. 5). A trend of increasing areal net methanogenesis rates
- 513 from March to September was observed in both years.

514 3.4.2 Hydrogenotrophic methanogenesis

- 515 Hydrogenotrophic methanogenesis activity determined by ¹⁴C-bicarbonate incubations of MUC cores
- 516 is shown in Fig. 2 and 3. In 2013, maximum activity ranged between 0.01-0.2 nmol cm⁻³ d⁻¹, while in
- 517 2014 maxima ranged only between 0.01 and 0.05 nmol cm⁻³ d⁻¹. In comparison, maximum
- 518 hydrogenotrophic methanogenesis was up to two orders of magnitude lower compared to net
- 519 methanogenesis. Only in March 2013 both activities reached a similar range.
- 520 Overall, hydrogenotrophic methanogenesis increased with depth in March, September, and
- 521 November 2013 and in March, June, and September 2014. In June 2013, activity decreased with
- depth, showing the highest rates in the upper 0-5 cmbsf and the lowest at the deepest sampleddepth.
- 524 Concomitant with integrated net methanogenesis, integrated hydrogenotrophic methanogenesis
- rates (0-25 cmbsf) were high in September 2013, with slightly higher rates in March 2013 (Fig. 5).
- 526 Lowest areal rates of hydrogenotrophic methanogenesis were seen in June of both years.
- 527 Hydrogenotrophic methanogenesis activity in the gravity core is shown in Fig. 4. Highest activity (~
- 528 0.7 nmol cm $^{-3}$ d $^{-1}$) was measured at 45 cmbsf and 138 cmbsf, followed by a decrease with increasing
- sediment depth reaching 0.01 nmol cm⁻³ d⁻¹ at the deepest sampled depth (345 cmbsf).
- 530 **3.4.3** Potential methanogenesis in manipulated experiments
- 531 Potential methanogenesis rates in manipulated experiments included either the addition of
- 532 inhibitors (molybdate for inhibition of sulfate reduction or BES for inhibition of methanogenesis) or
- 533 the addition of a non-competitive substrate (methanol). Control treatments were run with neither
- the addition of inhibitors nor the addition of methanol.
- 535 Controls. Potential methanogenesis activity in the control treatments was below 0.5 nmol cm⁻³ d⁻¹
- 536 from March 2014 to September 2014 (Fig. 6). Only in November 2013, control rates exceeded 0.5

- 537 nmol cm⁻³ d⁻¹ below 6 cmbsf. While rates increased with depth in November 2013 and June 2014,
- 538 they decreased with depth at the other two sampling months.
- 539 Molybdate. Peak potential methanogenesis rates in the molybdate treatments were found in the
- 540 uppermost sediment interval (0-1 cmbsf) at almost every sampling month with rates being 3-30
- times higher compared to the control treatments (< 0.5 nmol cm⁻³ d⁻¹). In November 2013, potential 541
- 542 methanogenesis showed two maxima (0-1 and 10-15 cmbsf). Highest measured rates were found in

September 2014 (~6 nmol cm⁻³ d⁻¹), followed by November 2013 (~5 nmol cm⁻³ d⁻¹). 543

- 544 BES. Profiles of potential methanogenesis in the BES treatments were similar to the controls mostly
- 545 in the lower range < 0.5 nmol cm⁻³ d⁻¹. Only in November 2013 rates exceeded 0.5 nmol cm⁻³ d⁻¹.
- 546 Rates increased with depth at all sampling months, except for September 2014, where highest rates 547 were found at the sediment surface (0-1 cmbsf).
- Methanol. At all sampling months, potential rates in the methanol treatments were three orders of 548
- 549 magnitude higher compared to the control treatments (< 0.5 nmol cm⁻³ d⁻¹). Except for November
- 550 2013, potential methanogenesis rates in the methanol treatments were highest in the upper 0-5
- 551 cmbsf and decreased with depth. In November 2013, highest rates were detected at the deepest 552 sampled depth (20-25 cmbsf).
- 553

560

554 3.4.4 Potential methanogenesis followed by ¹³C-methanol labeling

- The concentration of total methanol concentrations (labeled and unlabeled) in the sediment 555
- decreased sharply in the first 2 weeks from ~8 mM at day 1 to 0.5 mM at day 13 (Fig. 7). At day 17, 556 557 methanol was below the detection limit. In the first 2 weeks, residual methanol was enriched with
- ¹³C, reaching ~200 ‰ at day 13. 558
- Over the same time period, the methane content in the headspace increased from 2 ppmv at day 1 559
- to ~ 66,000 ppmv at day 17 and stayed around that value until the end of the total incubation time (until day 37) (Fig. 7). The carbon isotopic signature of methane ($\delta^{13}C_{CH4}$) showed a clear enrichment 561
- 562 of the heavier isotope ¹³C (Table 3) from day 9 to 17 (no methane was detectable at day 1). After day
- 17, $\delta^{13}C_{CH4}$ stayed around 13‰ until the end of the incubation. The content of CO₂ in the headspace 563
- 564 increased from ~8900 ppmv at day 1 to ~29,000 ppmv at day 20 and stayed around 30,000 ppmv
- 565 until the end of the incubation (Fig. 7). Please note, that the major part of CO₂ was dissolved in the
- 566 porewater, thus the CO_2 content in the headspace does not show the total CO_2 abundance in the
- 567 system. CO₂ in the headspace was enriched with ¹³C during the first 2 weeks (from -16.2 to -7.3 ‰)
- but then stayed around -11 ‰ until the end of the incubation. 568

569 3.5 Molecular analysis of benthic methanogens

- 570 In September 2014, additional samples were run during the methanol treatment (see Sect. 2.7.) for
- the detection of benthic methanogens via qPCR. The qPCR results are shown in Fig. 8. For a better 571

- 572 comparison, the microbial abundances are plotted together with the sediment methane
- 573 concentrations from the methanol treatment, from which the rate calculation for the methanol-
- 574 methanogenesis at 0-1 cmbsf was done (shown in Fig. 6).
- 575 Sediment methane concentrations increased over time revealing a slow increase in the first ~10 days,
- 576 followed by a steep increase between day 13 and day 20 and ending in a stationary phase.
- 577 A similar increase was seen in the abundance of total and methanogenic archaea. Total archaea
- 578 abundances increased sharply in the second week of the incubation reaching a maximum at day 16
- 579 (~5000 $*10^{6}$ copies g⁻¹) and stayed around 3000 $*10^{6}$ -4000 $*10^{6}$ copies g⁻¹ over the course of the
- 580 incubation. Similarly, methanogenic archaea, namely the order *Methanosarcinales* and within this
- 581 order the family Methanosarcinaceae, showed a sharp increase in the first 2 weeks as well with the
- highest abundances at day 16 (6* 10⁸ copies g⁻¹ and $^{1*10^6}$ copies g⁻¹, respectively). Until the end of
- 583 the incubation, the abundances of Methanosarcinales and Methanosarcinaceae decreased to about a
- 584 third of their maximum abundances ($^{2*10^8}$ copies g^{-1} and $^{-0.4*10^6}$ copies g^{-1} , respectively).

585 3.6 Statistical Analysis

- 586 The PCA of integrated SRZ methanogenesis (0-5 cmbsf) (Fig. 10) showed a positive correlation with
- 587 bottom water temperature (Fig. 10a), bottom water salinity (Fig. 10a), bottom water methane (Fig.
- 588 10b), surface sediment POC content (0-5 cmbsf, Fig. 10c), and surface sediment C/N (0-5 cmbsf, Fig.
- 10b). A negative correlation was found with bottom water oxygen concentration (Fig. 10b). No
- 590 correlation was found with bottom water chlorophyll.
- 591 The PCA of methanogenesis depth profiles showed positive correlations with sediment depth (Fig.
- 592 11a) and C/N (Fig. 11b), and showed negative correlations with POC (Fig. 11a).
- 593

594 **4. Discussion**

595 4.1 Methanogenesis in the sulfate-reducing zone

596 On the basis of the results presented in Fig. 2 and 3, it is evident that methanogenesis and sulfate 597 reduction were concurrently active in the sulfate reduction zone (0-30 cmbsf) at Boknis Eck. Even 598 though sulfate reduction activity was not directly determined, the decrease in sulfate concentrations with a concomitant increase in sulfide within the upper 30 cmbsf clearly indicated its presence (Fig. 2 599 600 and 3). Several previous studies confirmed the high activity of sulfate reduction in the surface 601 sediment of Eckernförde Bay, revealing rates up to 100-10,000 nmol cm⁻³ d⁻¹ in the upper 25 cmbsf 602 (Treude et al., 2005a; Bertics et al., 2013; Dale et al., 2013). Microbial fermentation of organic matter 603 was probably high in the organic-rich sediments of Eckernförde Bay (POC contents of around 5 %, Fig.

604 2 and 3), providing high substrate availability and variety for methanogenesis.

605							
606	The results of this study further identified methylotrophy to be a potentially important non-						
607	competitive methanogenic pathway in the sulfate-reducing zone. The pathway utilizes alternative						
608	substrates, such as methanol, to bypass competition with sulfate reducers for H_2 and acetate. A						
609	potential for methylotrophic methanogenesis within the sulfate-reducing zone was supported by the						
610	following observations:,						
611	1)	Hydrogenotrophic methanogenesis was up to two orders of magnitude lower compared to					
612		net methanogenesis, resulting in insufficient rates to explain the observed net					
613		methanogenesis in the upper 0-30 cmbsf (Fig. 2 and 3). This finding points towards the					
614		presence of alternative methanogenic processes in the sulfate reduction zone, such as					
615		methylotrophic methanogenesis.					
616	2)	Methanogenesis increased when sulfate reduction was inhibited by molybdate, confirming					
617		the inhibitory effect of sulfate reduction on methanogenesis with competitive substrates (H $_{\rm 2}$					
618		and acetate (Oremland & Polcin, 1982; King et al., 1983)) (Fig. 6), Consequently, usage of					
619		non-competitive substrates was preferred in sulfate reduction zone (especially in the upper					
620		0-1 cmbsf, Fig. 6). Accordingly, hydrogenotrophic methanogenesis increased at depths where					
621		sulfate was depleted and thus the competitive situation was relieved (Fig. 4).					
622	3)	3) The addition of BES did not result in the inhibition of methanogenesis, indicating the					
623		presence of unconventional methanogenic groups using non-competitive substrates (Fig. 7).					
624		The unsuccessful inhibition by BES can be explained either by incomplete inhibition or by the					
625		fact that the methanogens were insensitive to BES (Hoehler et al., 1994; Smith & Mah, 1981;					
626		Santoro & Konisky, 1987). The BES concentration applied in the present study (60 mM) has					
627		been shown to result in successful inhibition of methanogens in previous studies (Hoehler et					
628		al., 1994). Therefore, the presence of methanogens that are insensitive to BES is more likely.					
629		The insensitivity to BES in methanogens is explained by heritable changes in BES permeability					
630		or formation of BES-resistant enzymes (Smith & Mah, 1981; Santoro & Konisky, 1987). Such					
631		BES resistance was found in Methanosarcina mutants (Smith & Mah, 1981; Santoro &					
632		Konisky, 1987). This genus was successfully detected in our samples (for more details see					
633		point 5), and is known for mediating the methylotrophic pathway (Keltjens & Vogels, 1993),					
634		supporting our hypothesis on the utilization of non-competitive substrates by methanogens.					
635	4)	The addition of methanol to sulfate-rich sediments increased methanogenesis rates up to					
636		three orders of magnitude, confirming the potential of the methanogenic community to					
637		utilize non-competitive substrates especially in the 0-5 cmbsf sediment horizon (Fig. 6). At					
638		this sediment depth either the availability of non-competitive substrates, including					
639		methanol, was highest (derived from fresh organic matter), or the usage of non-competitive					

640		substrates was increased due to the high competitive situation as sulfate reduction is most
641		active in the 0-5 cmbsf layer (Treude et al., 2005a; Bertics et al., 2013). It should be noted
642		that even though methanogenesis rates were calculated assuming a linear increase in
643		methane concentration over the entire incubation to make a better comparison between
644		different treatments, the methanol treatments generally showed a delayed response in
645		methane development (Fig. 8, Supplement, Fig. S2). We suggest that this delayed response
646		was a reflection of cell growth by methanogens utilizing the surplus methanol. We are
647		therefore unable to decipher whether methanol plays a major role as a substrate in the
648		Eckernförde Bay sediments compared to possible alternatives, as its concentration is
649		relatively low in the natural setting (~1 μM between 0 and 25 cmbsf, June 2014 sampling, G
650		C. Zhuang unpubl. data). It is conceivable that other non-competitive substrates, such as
651		methylated sulfides (e.g., dimethyl sulfide or methanethiol), are more relevant for the
652		support of SRZ methanogenesis.
653	5)	Methylotrophic methanogens of the order Methanosarcinales were detected in the
654		methanol-treatment (Fig. 8), confirming the presence of methanogens that utilize non-
655		competitive substrates in the natural environment (Boone et al., 1993), (Fig. 8). The delay in
656		growth of Methanosarcinales moreover hints towards the predominant usage of other non-
657		competitive substrates over methanol (see also point 4).
658	6)	Stable isotope probing revealed highly 13 C-enriched methane produced from 13 C-labelled
659		methanol, furthermore confirming the potential of the methanogenic community to utilize
660		non-competitive substrates (Fig. 7). The production of both methane and CO_2 from methanol
661		has been shown previously in different strains of methylotrophic methanogens (Penger et al.,
662		2012). The fast conversion of methanol to methane and CO_2 (methanol was consumed
663		completely in 17 days) is hinting towards the presence of methylotrophic methanogens (e.g.
664		members of the family Methanosarcinaceae, which is known for the methylotrophic pathway
665		(Keltjens & Vogels, 1993)). Please note, however, that the storage of the cores (3.5 months)
666		prior to sampling could have led to shifts in the microbial community and thus might not
667		reflect in-situ conditions of the original microbial community in September 2014. The delay in
668		methane production also seen in the stable isotope experiment was, however, only slightly
669		different (methane developed earlier, between day 8 and 12, data not shown) from the non-
670		labeled methanol treatment (between day 10 to 16, Fig. S2), which leads us to the
671		assumption that the storage time at 1°C did not dramatically affect the methanogen
672		community. Similar, in a previous study with arctic sediments, addition of substrates had no
673		stimulatory effect on the rate of methanogenesis or on the methanogen community
674		structure at low temperatures (5°C, (Blake et al., 2015)).

675 4.2 Environmental control of methanogenesis in the sulfate reduction zone

- 676 SRZ methanogenesis in Eckernförde Bay sediments showed variations throughout the sampling
- 677 period, which may be influenced by variable environmental factors such as temperature, salinity,
- 678 oxygen, and organic carbon. In the following, we will discuss the potential impact of those factors on
- the magnitude and distribution of SRZ methanogenesis.

680 4.2.1 Temperature

681 During the sampling period, bottom water temperatures increased over the course of the year from 682 late winter (March, 3-4 °C) to autumn (November, 12°C, Fig. 2 and 3). The PCA revealed a positive 683 correlation between bottom water temperature and integrated SRZ methanogenesis (0-5 cmbsf). A 684 temperature experiment conducted with sediment from ~75 cmbsf in September 2014 within a 685 parallel study revealed a mesophilic temperature optimum of methanogenesis (20 °C, data not 686 shown). Whether methanogenesis in the sulfate reduction zone (0-30 cm) has the same physiology 687 remains speculative. However, AOM organisms, which are closely related to methanogens (Knittel & 688 Boetius, 2009), studied in the sulfate reduction zone from the same site were confirmed to have a mesophilic physiology, too (Treude et al. 2005). The sum of these aspects lead us to the conceivable 689 690 conclusion that SRZ methanogenesis activity in the Eckernförde Bay is positively impacted by 691 temperature increases. Such a correlation between benthic methanogenesis and temperature has 692 been found in several previous studies from different environments ((Sansone & Martens, 1981; Crill 693 & Martens, 1983; Martens & Klump, 1984).

694

695 4.2.2 Salinity and oxygen

696 From March 2013 to November 2013, and from March 2014 to September 2014, salinity increased in the bottom-near water (25 m) from 19 to 23 PSU and from 22 to 25 PSU (Fig. 2 and 3), respectively, 697 due the pronounced summer stratification in the water column between saline North Sea water and 698 less saline Baltic Sea water (Bange et al., 2011). The PCA detected a positive correlation between 699 700 integrated SRZ methanogenesis (0-5 cmbsf) and salinity in the bottom-near water (Fig. 10a). This 701 correlation can hardly be explained by salinity alone, as methanogens feature a broad salinity range 702 from freshwater to hypersaline (Zinder, 1993). More likely, salinity serves as an indicator of water-703 column stratification, which is often correlated with low O2 concentrations in the Eckernförde Bay 704 (Fig. S3, Bange et al., 2011; Bertics et al., 2013). Methanogenesis is sensitive to O₂ (Oremland, 1988; 705 Zinder, 1993), and hence conditions might be more favorable during hypoxic or anoxic events, 706 particular in the sediment closest to the sediment-water interface, but potentially also in deeper 707 sediment layers due to the absence of bioturbating and bioirrigating infauna (Dale et al., 2013; 708 Bertics et al., 2013), which could introduce O₂ beyond diffusive transport. Accordingly, the PCA

709 revealed a negative correlation between O₂ concentration close to the seafloor and SRZ

- 710 methanogenesis.
- 711

712 4.2.4 Particulate organic carbon

713 The supply of particulate organic carbon (POC) is one of the most important factors controlling

714 benthic heterotrophic processes, as it determines substrate availability and variety (Jørgensen,

715 2006). In Eckernförde Bay, the organic material reaching the seafloor originates mainly from

716 phytoplankton blooms in spring, summer and autumn (Bange et al., 2011). It has been estimated that

717 >50 % in spring (February/March), <25 % in summer (July/August) and >75 % in autumn

718 (September/October) of these blooms is reaching the seafloor (Smetacek et al., 1984), resulting in a

719 overall high organic carbon content of the sediment (5 wt %), which leads to high benthic microbial

720 degradation rates including sulfate reduction and methanogenesis (Whiticar, 2002; Treude et al.,

721 2005a; Bertics et al., 2013). Previous studies revealed that high organic matter availability can relieve

722 competition between sulfate reducers and methanogens in sulfate-containing, marine sediments

723 (Oremland et al., 1982; Holmer & Kristensen, 1994; Treude et al., 2009; Maltby et al., 2016).

To determine the effect of POC concentration and C/N ratio (the latter as a negative indicator for the

725 freshness of POC) on SRZ methanogenesis, two PCAs were conducted with a) the focus on the upper

726 0-5 cmbsf, which is directly influenced by freshly sedimented organic material from the water column

(Fig. 10), and b) the focus on the depth profiles throughout the sediment cores (up to 30 cmbsf) (Fig.11).

729 a) Effect of POC and C/N ratio in the upper 0-5 cmbsf

730 For the upper 0-5 cmbsf in the sediment, a positive correlation was found between SRZ methanogenesis (integrated) and POC content (averaged) (Fig. 10c), indicating that POC content is an 731 important controlling factor for methanogenesis in this layer. In support, highest bottom-near water 732 733 chlorophyll concentrations coincided with highest bottom-near water methane concentrations and 734 high integrated SRZ methanogenesis (0-5 cmbsf) in September 2013, probably as a result of the 735 sedimentation of the summer phytoplankton bloom (Fig. 9). Indeed, the PCA revealed a positive 736 correlation between integrated SRZ methanogenesis rates and bottom-near water methane concentrations (Fig. 10b), when viewed over all investigated months. However, no correlation was 737 738 found between bottom water chlorophyll and integrated SRZ methanogenesis rates (Fig. 10). As seen in Fig. 9, bottom-near high chlorophyll concentrations did not coincide with high bottom-near 739 740 methane concentration in June/September 2014. We explain this result by a time lag between 741 primary production in the water column and the export of the produced organic material to the 742 seafloor, which was probably even more delayed during stratification. Such a delay was observed in a 743 previous study (Bange et al., 2010), revealing enhanced water methane concentration close to the

744 seafloor approximately one month after the chlorophyll maximum. The C/N ratio (averaged over 0-5 745 cmbsf) also showed no correlation with integrated methanogenesis from the same depth layer (0-5 746 cmbsf), which is surprising as we expected that a higher C/N ratio, indicative for less labile organic 747 carbon, should have a negative effect on non-competitive methanogenesis. However, methanogens are not able to directly use most of the labile organic matter due their inability to process large 748 749 molecules (more than two C-C bondings) (Zinder, 1993). Methanogens are dependent on other 750 microbial groups to degrade large organic compounds (e.g. amino acids) for them (Zinder, 1993). 751 Because of this substrate speciation and dependence, a delay between the sedimentation of fresh, 752 labile organic matter and the increase in methanogenesis can be expected, which would not be 753 captured by the applied PCA.

754 b) Effect of POC and C/N ratio over 0-30 cmbsf

755 In the PCA for the sediment profiles from the sulfate reduction zone (0-30 cmbsf), POC showed a 756 negative correlation with methanogenesis and sediment depth, while C/N ratio showed a positive 757 correlation with methanogenesis and sediment depth (Fig 11.). Given that POC remained basically 758 unchanged over the top 30 cmbsf, with the exemption of the topmost sediment layer, its negative 759 correlation with methanogenesis is probably solely explained by the increase of methanogenesis 760 with sediment depth, and can therefore be excluded as a major controlling factor. As sulfate in this 761 zone was likely never depleted to levels that are critically limiting sulfate reduction (lowest 762 concentration 1300 µM, compare e.g. with Treude et al., 2014) we do not expect a significant change 763 in the competition between methanogens and sulfate reducers. It is therefore more likely that the 764 progressive degradation of labile POC into dissolvable methanogenic substrates over depth and time 765 had a positive impact on methanogenesis. The C/N ratio indicates such a trend as the labile fraction of POC decreased with depth. 766

767 4.3 Relevance of methanogenesis in the sulfate reduction zone of Eckernförde Bay sediments

The time series station Boknis Eck in Eckernförde Bay is known for being a methane source to the
atmosphere throughout the year due to supersaturated waters, which result from significant benthic
methanogenesis and emission (Bange et al., 2010). The benthic methane formation is thought to take
place mainly in sediments below the SMTZ (Treude et al., 2005a; Whiticar, 2002).
In the present study, we show that SRZ methanogenesis within the sulfate zone is present despite

- sulfate concentrations > 1 mM, a limit above which methanogenesis has been thought to be
- negligible (Alperin et al., 1994; Hoehler et al., 1994; Burdige, 2006), and thus could contribute to
- benthic methane emissions. In support of this hypothesis, high dissolved methane concentration in
- the water column occurred with concomitant high SRZ methanogenesis activity (Fig. 9). However,
- 777 whether the observed water-column methane originated from SRZ methanogenesis or from gas
- 778 ebullition caused by methanogenesis below the SMTZ, or a mixture of both, remains speculative.

779 How much of the methane produced in the surface sediment is ultimately emitted into the water 780 column depends on the rate of methane consumption, i.e., aerobic and anaerobic oxidation of 781 methane in the sediment (Knittel & Boetius, 2009) (Fig. 1). In organic-rich sediments, such as in the 782 presented study, the oxygenated sediment layer is often only mm-thick, due to the high O₂ demand of microorganisms during organic matter degradation (Jørgensen, 2006; Preisler et al., 2007). Thus, 783 784 the anaerobic oxidation of methane (AOM) might play a more important role for methane 785 consumption in the studied Eckernförde Bay sediments. In an earlier study from this site, AOM 786 activity was detected throughout the top 0-25 cmbsf, which included zones that were well above the 787 actual SMTZ (Treude et al., 2005a). But the authors concluded that methane oxidation was 788 completely fueled by methanogenesis from below sulfate penetration, as integrated AOM rates (0.8-789 1.5 mmol m⁻² d⁻¹) were in the same range as the predicted methane flux (0.66-1.88 mmol m⁻² d⁻¹) into 790 the SMT7. 791 Together with the data set presented here we postulate that AOM above the SMTZ (0.8 mmol m⁻² d⁻¹, 792 Treude et al., (2005a) could be partially or entirely fueled by SRZ methanogenesis. A similar close 793 coupling between methane oxidation and methanogenesis in the absence of definite methane profiles was recently proposed from isotopic labeling experiments with sediments from the sulfate 794 reduction zone of the close-by Aarhus Bay, Denmark (Xiao et al., 2017). It is therefore likely that such 795 796 a cryptic methane cycling also occurs in the sulfate reduction zone of sediments in the Eckernförde 797 Bay. If, in an extreme scenario, SRZ methanogenesis would represent the only methane source for AOM above the SMTZ, then maximum SRZ methanogenesis could be in the order of 1.6 mmol m $^{-2}$ d $^{-1}$ 798 799 (1.5 mmol m⁻² d⁻¹ AOM + 0.09 mmol m⁻² d⁻¹ net SRZ methanogenesis). 800 Even though the contribution of SRZ methanogenesis to AOM above the SMTZ remains speculative, it 801 leads to the assumption that SRZ methanogenesis could play a much bigger role for benthic carbon 802 cycling in the Eckernförde Bay than previously thought. Whether SRZ methanogenesis at Eckernförde

- 803 Bay has the potential for the direct emission of methane into the water column goes beyond the
- scope of this study and should be tested in the future.

805 5. Summary

- The present study demonstrated that methanogenesis and sulfate reduction were concurrently active within the sulfate-reducing zone in sediments at Boknis Eck (Eckernförde Bay, SW Baltic Sea). The observed methanogenesis was probably based on non-competitive substrates due to the competition with sulfate reducers for the substrates H₂ and acetate. Accordingly, members of the family *Methanosarcinaceae*, which are known for methylotrophic methanogenesis, were found in the sulfate reduction zone of the sediments and are likely to be responsible for the observed methanogenesis with the potential use of non-competitive substrates such as methanol,
- 813 methylamines or methylated sulfides.

- 814 Potential environmental factors controlling SRZ methanogenesis are POC content, C/N ratio, oxygen,
- and temperature, resulting in highest methanogenesis activity during the warm, stratified, and
- 816 hypoxic months after the late summer phytoplankton blooms.
- 817 This study provides new insights into the presence and seasonality of SRZ methanogenesis in coastal
- 818 sediments, and was able to demonstrate that the process could play an important role for the
- 819 methane budget and carbon cycling of Eckernförde Bay sediments, e.g., by directly fueling AOM
- 820 above the SMTZ.
- 821

822 Author Contribution

- 823 J.M. and T.T. designed the experiments. J.M. carried out all experiments. H.W.B. coordinated
- 824 measurements of water column methane and chlorophyll. C.R.L. and M.A.F. conducted molecular
- 825 analysis. M.S. coordinated 13C-Isotope measurements. J.M. prepared the manuscript with
- 826 contributions from all co-authors.

827 Data Availability

- 828 Research data for the present study can be accessed via the public data repository PANGEA
- 829 (doi:10.1594/PANGAEA.873185).

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846 References

847 848 849	Abegg, F. & Anderson, A.L. (1997). The acoustic turbid layer in muddy sediments of Eckernfoerde Bay , Western Baltic : methane concentration , saturation and bubble characteristics. <i>Marine Geology</i> . 137. pp. 137–147.
850	Alperin, M.J., Albert, D.B. & Martens, C.S. (1994). Seasonal variations in production and consumption
851	rates of dissolved organic carbon in an organic-rich coastal sediment. <i>Geochimica et</i>
852	<i>Cosmochimica Acta</i> . 58 (22). pp. 4909–4930.
853	Bakker, D.E., Bange, H.W., Gruber, N., Johannessen, T., Upstill-Goddard, R.C., Borges, A.V., Delille, B.,
854	Löscher, C.R., Naqvi, S.W.A., Omar, A.M. & Santana-Casiano-J.M. (2014). Air-sea interactions of
855	natural long-lived greenhouse gases (CO2, N2O, CH4) in a changing climate. In: P. S. Liss & M. T.
856	Johnson (eds.). Ocean-Atmosphere Interactions of Gases and Particles. Heidelberg: Springer-
857	Verlag, pp. 113–169.
858	Balzer, W., Pollehne, F. & Erlenkeuser, H. (1986). Cycling of Organic Carbon in a Marine Coastal
859	System. In: P. G. Sly (ed.). <i>Sediments and Water Interactions</i> . New York, NY: Springer New York,
860	pp. 325–330.
861	Bange, H.W., Bartell, U.H., Rapsomanikis, S. & Andreae, M.O. (1994). Methane in the Baltic and North
862	Seas and a reassessment of the marine emissions of methane. <i>Global Biogeochemical Cycles</i> . 8
863	(4). pp. 465–480.
864	Bange, H.W., Bergmann, K., Hansen, H.P., Kock, A., Koppe, R., Malien, F. & Ostrau, C. (2010).
865	Dissolved methane during hypoxic events at the Boknis Eck time series station (Eckernförde
866	Bay , SW Baltic Sea). <i>Biogeosciences</i> . 7. pp. 1279–1284.
867	Bange, H.W., Hansen, H.P., Malien, F., Laß, K., Karstensen, J., Petereit, C., Friedrichs, G. & Dale, A.
868	(2011). Boknis Eck Time Series Station (SW Baltic Sea): Measurements from 1957 to 2010.
869	LOICZ-Affiliated Activities. Inprint 20. pp. 16–22.
870	Bertics, V.J., Löscher, C.R., Salonen, I., Dale, A.W., Gier, J., Schmitz, R.A. & Treude, T. (2013).
871	Occurrence of benthic microbial nitrogen fixation coupled to sulfate reduction in the seasonally
872	hypoxic Eckernförde Bay, Baltic Sea. <i>Biogeosciences</i> . 10 (3). pp. 1243–1258.
873	Blake, L.I., Tveit, A., Øvreås, L., Head, I.M. & Gray, N.D. (2015). Response of Methanogens in Arctic
874	Sediments to Temperature and Methanogenic Substrate Availability. <i>Planet. Space Sci.</i> 10 (6).
875	pp. 1–18.
876	Buckley, D.H., Baumgartner, L.K. & Visscher, P.T. (2008). Vertical distribution of methane metabolism
877	in microbial mats of the Great Sippewissett Salt Marsh. <i>Environmental microbiology</i> . 10 (4). pp.
878	967–77.
879 880	Burdige, D.J. (2006). <i>Geochemistry of Marine Sediments</i> . New Jersey, U.S.A.: Princeton University Press.
881 882	Cicerone, R.J. & Oremland, R.S. (1988). Biogeochemical aspects of atmospheric methane. <i>Global Biogeochemical Cycles</i> . 2 (4). pp. 299–327.
883 884	Crill, P. & Martens, C. (1983). Spatial and temporal fluctuations of methane production in anoxic coastal marine sediments. <i>Limnology and Oceanography</i> . 28. pp. 1117–1130.
885 886	Crill, P.M. & Martens, C.S. (1986). Methane production from bicarbonate and acetate in an anoxic marine sediment. <i>Geochimica et Cosmochimica Acta</i> . 50. pp. 2089–2097.
887 888	Dale, a. W., Bertics, V.J., Treude, T., Sommer, S. & Wallmann, K. (2013). Modeling benthic-pelagic

nutrient exchange processes and porewater distributions in a seasonally hypoxic sediment:
evidence for massive phosphate release by Beggiatoa? *Biogeosciences*. 10 (2). pp. 629–651.

890 Denman, K.L., Brasseur, G., Chidthaisong, A., Ciais, P., Cox, P.M., Dickinson, R.E., Hauglustaine, D.,

- 891 Heinze, C., Holland, E., Jacob, D., Lohmann, U., Ramachandran, S., da Silva Dias, P.L., Wofsy, S.C.
- 892 & Zhang, X. (2007). Couplings Between Changes in the Climate System and Biogeochemistry. In:
- S. Solomon, D. Qin, M. Manning, Z. Chen, M. Marquis, K. B. Averyt, M. Tignor, & H. L. Miller
 (eds.). Climate Change 2007: The Physical Science Basis. Contribution of Wokring Group I to th
- (eds.). Climate Change 2007: The Physical Science Basis. Contribution of Wokring Group I to the
 Fourth Assessment Report of the Intergovernmental Panel on Climate Change. Cambridge,
 United Kingdom and New York, NY, USA: Cambridge University Press.
- 897 EPA (2010). Methane and nitrous oxide emissions from natural sources. Washington, DC, USA.
- Ferdelman, T.G., Lee, C., Pantoja, S., Harder, J., Bebout, B.M. & Fossing, H. (1997). Sulfate reduction
 and methanogenesis in a Thioploca-dominated sediment off the coast of Chile. *Geochimica et Cosmochimica Acta*. 61 (15). pp. 3065–3079.
- 901 Gier, J., Sommer, S., Löscher, C.R., Dale, A.W., Schmitz, R.A. & Treude, T. (2016). Nitrogen fixation in
 902 sediments along a depth transect through the Peruvian oxygen minimum zone. *Biogeosciences*.
 903 13 (14). pp. 4065–4080.
- 904 Grasshoff, K., Ehrhardt, M. & Kremmling, K. (1999). *Methods of Seawater Analysis*. Weinheim: Verlag 905 Chemie.
- Hansen, H.-P., Giesenhagen, H.C. & Behrends, G. (1999). Seasonal and long-term control of bottom water oxygen deficiency in a stratified shallow-water coastal system. *ICES Journal of Marine Science*. 56. pp. 65–71.
- Hartmann, D.L., Klein Tank, A.M.G., Rusticucci, M., Alexander, L.V., Brönnimann, S., Charabi, Y.,
 Dentener, F.J., Dlugokencky, D.R., Easterling, D.R., Kaplan, A., Soden, B.J., Thorne, P.W., Wild,
 M. & Zhai, P.M. (2013). Observations: Atmosphere and Surface. In: *Climate Change 2013: The pHysical Science Basis. Contribution Group I to the Fifth Assessment Report of the*Intergovernmental Panel on Climate Change. United Kingdom and New York, NY, USA:
- 914 Cambridge University Press.
- Hoehler, T.M., Alperin, M.J., Albert, D.B. & Martens, C.S. (1994). Field and laboratory studies of
 methane oxidation in an anoxic marine sediment: Evidence for a methanogen-sulfate reducer
 consortium. *Global Biogeochemical Cycles*. 8 (4). pp. 451–463.
- 918Holmer, M. & Kristensen, E. (1994). Coexistence of sulfate reduction and methane production in an919organic-rich sediment. Marine Ecology Progress Series. 107. pp. 177–184.
- Jackson, D.R., Williams, K.L., Wever, T.F., Friedrichs, C.T. & Wright, L.D. (1998). Sonar evidence for
 methane ebullition in Eckernforde Bay. *Continental Shelf Research*. 18. pp. 1893–1915.
- Jørgensen, B.B. (2006). Bacteria and marine Biogeochemistry. In: H. D. Schulz & M. Zabel (eds.).
 Marine Geochemistry. Berlin/Heidelberg: Springer-Verlag, pp. 173–207.
- Jørgensen, B.B. & Parkes, R.J. (2010). Role of sulfate reduction and methane production by organic
 carbon degradation in eutrophic fjord sediments (Limfjorden, Denmark). *Limnology and Oceanography*. 55 (3). pp. 1338–1352.
- Keltjens, J.T. & Vogels, G.D. (1993). Conversion of methanol and methylamines to methane and
 carbon dioxide. In: J. G. Ferry (ed.). *Methanogenesis: Ecology, Physiology, Biochemistry & Genetics*. Chapman & Hall, pp. 253–303.
- King, G.M., Klug, M.J. & Lovley, D.R. (1983). Metabolism of acetate, methanol, and methylated
 amines in intertidal sediments of lowes cove, maine. *Applied and environmental microbiology*.
 45 (6). pp. 1848–1853.
- Knittel, K. & Boetius, A. (2009). Anaerobic oxidation of methane: progress with an unknown process.
 Annual review of microbiology. 63. pp. 311–34.
- Lennartz, S.T., Lehmann, A., Herrford, J., Malien, F., Hansen, H.-P., Biester, H. & Bange, H.W. (2014).
 Long-term trends at the Boknis Eck time series station (Baltic Sea), 1957–2013: does climate
 change counteract the decline in eutrophication? *Biogeosciences*. 11 (22). pp. 6323–6339.

- Maltby, J., Sommer, S., Dale, A.W. & Treude, T. (2016). Microbial methanogenesis in the sulfate reducing zone of surface sediments traversing the Peruvian margin. *Biogeosciences*. 13. pp.
 283–299.
- Martens, C.S., Albert, D.B. & Alperin, M.J. (1998). Biogeochemical processes controlling methane in
 gassy coastal sediments---Part 1. A model coupling organic matter flux to gas production ,
 oxidation and transport. *Continental Shelf Research*. 18. pp. 14–15.
- Martens, C.S. & Klump, J. V (1984). Biogeochemical cycling in an organic-rich coastal marine basin 4.
 An organic carbon budget for sediments dominated by sulfate reduction and methanogenesis.
 Geochimica et Cosmochimica Acta. 48. pp. 1987–2004.
- Naqvi, S.W. a., Bange, H.W., Farías, L., Monteiro, P.M.S., Scranton, M.I. & Zhang, J. (2010). Marine
 hypoxia/anoxia as a source of CH4 and N2O. *Biogeosciences*. 7 (7). pp. 2159–2190.
- 949 Oremland, R.S. (1988). Biogeochemistry of methanogenic bacteria. In: A. J. B. Zehnder (ed.). *Biology* 950 of Anaerobic Microorganisms. New York: J. Wiley & Sons, pp. 641–705.
- 951 Oremland, R.S. & Capone, D.G. (1988). Use of specific inhibitors in biogeochemistry and microbial
 952 ecology. In: K. C. Marshall (ed.). Advances in Microbial Ecology. Advances in Microbial Ecology.
 953 Boston, MA: Springer US, pp. 285–383.
- 954 Oremland, R.S., Marsh, L.M. & Polcin, S. (1982). Methane production and simultanous sulfate
 955 reduction in anoxic,salt-marsh sediments. *Nature*. 286. pp. 143–145.
- Oremland, R.S. & Polcin, S. (1982). Methanogenesis and Sulfate Reduction : Competitive and
 Noncompetitive Substrates in Estuarine Sediments. *Applied and Environmental Microbiology*. 44
 (6). pp. 1270–1276.
- Orsi, T.H., Werner, F., Milkert, D., Anderson, a. L. & Bryant, W.R. (1996). Environmental overview of
 Eckernförde Bay, northern Germany. *Geo-Marine Letters*. 16 (3). pp. 140–147.
- Penger, J., Conrad, R. & Blaser, M. (2012). Stable carbon isotope fractionation by methylotrophic
 methanogenic archaea. *Applied and environmental microbiology*. 78 (21). pp. 7596–602.
- Pimenov, N., Davidova, I., Belyaev, S., Lein, A. & Ivanov, M. (1993). Microbiological processes in
 marine sediments in the Zaire River Delta and the Benguela upwelling region. *Geomicrobiology Journal*. 11 (3–4). pp. 157–174.
- Preisler, A., de Beer, D., Lichtschlag, A., Lavik, G., Boetius, A. & Jørgensen, B.B. (2007). Biological and
 chemical sulfide oxidation in a Beggiatoa inhabited marine sediment. *The ISME journal*. 1 (4).
 pp. 341–353.
- 969 Reeburgh, W. (2007). Oceanic methane biogeochemistry. Chemical Reviews. pp. 486–513.
- Sansone, F.J. & Martens, C.S. (1981). Methane Production from Acetate and Associated Methane
 Fluxes from Anoxic Coastal Sediments. *Science*. 211 (4483). pp. 707–709.
- Santoro, N. & Konisky, J. (1987). Characterization of bromoethanesulfonate-resistant mutants of
 Methanococcus voltae: Evidence of a coenzyme M transport system. *Journal of Bacteriology*.
 169 (2). pp. 660–665.
- Schlüter, M., Sauter, E., Hansen, H.-P. & Suess, E. (2000). Seasonal variations of bioirrigation in
 coastal sediments: modelling of field data. *Geochimica et Cosmochimica Acta*. 64 (5). pp. 821–
 834.
- Seeberg-Elverfeldt, J., Schluter, M., Feseker, T. & Kolling, M. (2005). Rhizon sampling of porewaters
 near the sediment-water interface of aquatic systems. *Limnology and Oceanography-Methods*.
 3. pp. 361–371.
- Smetacek, V. (1985). The Annual Cycle of Kiel Bight Plankton: A Long-Term Analysis. *Estuaries*. 8
 (June). pp. 145–157.
- 983 Smetacek, V., von Bodungen, B., Knoppers, B., Peinert, R., Pollehne, F., Stegmann, P. & Zeitzschel, B.

- 984 (1984). Seasonal stages characterizing the annual cycle of an inshore pelagic system. *Rapports* 985 *et Proces-Verbaux des Reunions Conseil International pour l'Exploration de la Mer.* 186. pp.
 986 126–135.
- Smith, M.R. & Mah, R. a. (1981). 2-Bromoethanesulfonate: A selective agent for isolating
 resistantMethanosarcina mutants. *Current Microbiology*. 6 (5). pp. 321–326.
- Steinle, L., Maltby, J., Treude, T., Kock, A., Bange, H.W., Engbersen, N., Zopfi, J., Lehmann, M.F. &
 Niemann, H. (2017). Effects of low oxygen concentrations on aerobic methane oxidation in
 seasonally hypoxic coastal waters. *Biogeosciences*. 14 (6). pp. 1631–1645.
- Thießen, O., Schmidt, M., Theilen, F., Schmitt, M. & Klein, G. (2006). Methane formation and
 distribution of acoustic turbidity in organic-rich surface sediments in the Arkona Basin, Baltic
 Sea. *Continental Shelf Research*. 26 (19). pp. 2469–2483.
- Treude, T., Krause, S., Maltby, J., Dale, A.W., Coffin, R. & Hamdan, L.J. (2014). Sulfate reduction and
 methane oxidation activity below the sulfate-methane transition zone in Alaskan Beaufort Sea
 continental margin sediments: Implications for deep sulfur cycling. *Geochimica et Cosmochimica Acta*. 144. pp. 217–237.
- 999 Treude, T., Krüger, M., Boetius, A. & Jørgensen, B.B. (2005a). Environmental control on anaerobic
 1000 oxidation of methane in the gassy sediments of Eckernförde Bay (German Baltic). *Limnology* 1001 *and Oceanography*. 50 (6). pp. 1771–1786.
- Treude, T., Niggemann, J., Kallmeyer, J., Wintersteller, P., Schubert, C.J., Boetius, A. & Jørgensen, B.B.
 (2005b). Anaerobic oxidation of methane and sulfate reduction along the Chilean continental
 margin. *Geochimica et Cosmochimica Acta*. 69 (11). pp. 2767–2779.
- Treude, T., Smith, C.R., Wenzhöfer, F., Carney, E., Bernardino, A.F., Hannides, A.K., Krgüer, M. &
 Boetius, A. (2009). Biogeochemistry of a deep-sea whale fall: Sulfate reduction, sulfide efflux
 and methanogenesis. *Marine Ecology Progress Series*. 382. pp. 1–21.
- Welschmeyer, N.A. (1994). Fluorometric analysis of chlorophyll a in the presence of chlorophyll b and
 pheopigments. *Limnology and Oceanography*. 39 (8). pp. 1985–1992.
- Wever, T.F., Abegg, F., Fiedler, H.M., Fechner, G. & Stender, I.H. (1998). Shallow gas in the muddy
 sediments of Eckernförde Bay, Germany. *Continental Shelf Research*. 18. pp. 1715–1739.
- 1012 Wever, T.F. & Fiedler, H.M. (1995). Variability of acoustic turbidity in Eckernförde Bay (southwest
 1013 Baltic Sea) related to the annual temperature cycle. *Marine Geology*. 125. pp. 21–27.
- Whiticar, M.J. (2002). Diagenetic relationships of methanogenesis, nutrients, acoustic turbidity,
 pockmarks and freshwater seepages in Eckernförde Bay. *Marine Geology*. 182. pp. 29–53.
- Widdel, F. & Bak, F. (1992). Gram-Negative Mesophilic Sulfate-Reducing Bacteria. In: A. Balows, H. G.
 Trüper, M. Dworkin, W. Harder, & K.-H. Schleifer (eds.). *The Prokaryotes*. New York, NY:
 Springer New York, pp. 3352–3378.
- Wuebbles, D.J. & Hayhoe, K. (2002). Atmospheric methane and global change. *Earth-Science Reviews*.
 57 (3–4). pp. 177–210.
- 1021 Xiao, K.Q., Beulig, F., Kjeldsen, K.U., Jørgensen, B.B. & Risgaard-Petersen, N. (2017). Concurrent
 1022 methane production and oxidation in surface sediment from Aarhus Bay, Denmark. *Frontiers in* 1023 *Microbiology*. pp. 1–12.
- 1024 Zinder, S.H. (1993). Physiological ecology of methanogens. In: J. G. Ferry (ed.). *Methanogenesis*. New
 1025 York, NY: Chapman & Hall, pp. 128–206.
- 1026

1027 Figure Captions

1028 **Figure 1:** Overview of processes relevant for benthic methane production, consumption, and

- 1029 emission in the Eckernförde Bay. The thickness of arrows for emissions and coupling between surface
- 1030 processes indicates the strength of methane supply. Note that this figure combines existing
- 1031 knowledge with results from the present study. See discussion for more details.
- 1032 **Figure 2:** Parameters measured in the water column and sediment in the Eckernförde Bay at each
- 1033 sampling month in the year 2013. Net methanogenesis (MG) and hydrogenotrophic (hydr.)
- 1034 methanogenesis rates are shown in triplicates with mean (solid line).
- 1035 Figure 3: Parameters measured in the water column and sediment in the Eckernförde Bay at each
- sampling month in the year 2014. Net methanogenesis (MG) and hydrogenotrophic (hydr.)
- 1037 methanogenesis rates are shown in triplicates with mean (solid line).
- 1038 Figure 4: Parameters measured in the sediment gravity core taken in the Eckernförde Bay in
- September 2013. Hydrogenotrophic (hydr.) methanogenesis rates are shown in triplicates with mean(solid line).
- 1041 Figure 5: Integrated net methanogenesis (MG) rates (determined by net methane production) and
- 1042 hydrogenotrophic MG rates (determined by radiotracer incubation) in surface sediments (0-25
- 1043 cmbsf) of Eckernförde Bay for different sampled time points.
- 1044Figure 6: Potential methanogenesis rates versus sediment depth in sediment sampled in November10452013, March 2014, June 2014 and September 2014. Presented are four different types of incubations
- 1046 (treatments): Control (blue symbols) is describing the treatment with sediment plus artificial
- 1047 seawater containing natural salinity (24 PSU) and sulfate concentrations (17 mM), molybdate (green
- 1048 symbols) is the treatment with addition of molybdate (22 mM), BES (purple symbols) is the
- treatment with 60 mM BES addition, and *methanol* (red symbols) is the treatment with addition of 10
 mM methanol. Shown are triplicates per depth interval and the mean as a solid line. Please note the
 different x-axis for the methanol treatment (red).
- Figure 7: Development of headspace gas content and isotope composition of methane (CH₄) and
 carbon dioxide (CO₂), and porewater methanol (CH₃OH) concentration and isotope composition
 during the 13C-labeling experiment (with sediment from the 0-2 cmbsf horizon in September 2014)
 with addition of ¹³C-enriched methanol (¹³C:¹²C = 1:1000). *Figure above:* Concentrations of porewater
 methanol (CH₃OH) and headspace content of methane (CH₄) and carbon dioxide (CO₂) over time. *Figure below:* Isotope composition of porewater CH₃OH, headspace CH₄, and headspace CO₂ over
 time. Shown are means (from triplicates) with standard deviation.

Figure 8: Sediment methane concentrations (with sediment from the 0-1 cmbsf in September 2014)
over time in the treatment with addition of methanol (10 mM) are shown above. Shown are triplicate
values per measurement. DNA copies of *Archaea, Methanosarcinales* and *Methanosarcinaceae* are
shown below in duplicates per measurement. Please note the secondary y-axis for *Methanosarcinales* and *Methanosarcinaceae*. More data are available for methane (determined in
the gas headspace) than from DNA samples (taken from the sediment) as sample volume for
molecular analyzes was limited.

Figure 9: Temporal development of integrated net surface methanogenesis (0-5 cmbsf) in the
sediment and chlorophyll (green) and methane concentrations (orange) in the bottom water (25 m).
Methanogenesis (MG) rates and methane concentrations are shown in means (from triplicates) with
standard deviation.

1070 Figure 10: Principle component analysis (PCA) from three different angles of integrated surface 1071 methanogenesis (0-5 cmbsf) and surface particulate organic carbon averaged over 0-5 cmbsf (surface 1072 sediment POC), surface C/N ratio averaged over 0-5 cmbsf (surface sediment C/N), bottom water salinity, bottom water temperature (T), bottom water methane (CH_4), bottom water oxygen (O_2), and 1073 1074 bottom water chlorophyll. Data were transformed into ranks before analysis. a) Correlation biplot of 1075 principle components 1 and 2, b) correlation biplot of principle components 1 and 3, c) correlation 1076 biplot of principle components 2 and 3. Correlation biplots are shown in a multidimensional space 1077 with parameters shown as green lines and samples shown as black dots. Parameters pointing into the same direction are positively related; parameters pointing in the opposite direction are 1078 1079 negatively related.

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Figure 11: Principle component analysis (PCA) from two different angles of net methanogenesis
depth profiles and sampling month (Month), sediment depth, depth profiles of particulate organic
carbon (POC) and C/N ratio (C/N). Data was transformed into ranks before analysis. a) Correlation
biplot of principle components 1 and 2, b) correlation biplot of principle components 1 and 3.
Correlation biplots are shown in a multidimensional space with parameters shown as green lines and
samples shown as black dots. Parameters pointing into the same direction are positively related;
parameters pointing in the opposite direction are negatively related.

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Table 1: Sampling months with bottom water (~ 2 m above seafloor) temperature (Temp.), dissolved

Sampling Month	Date	Instrument	Temp. (°C)	O₂ (µM)	CH₄ (nM)	Type of Analysis
March 2013	13.03.2013	CTD	3	340	30	WC
		MUC				All
Juni 2013	27.06.2013	CTD	6	94	125	WC
		MUC				All
September 2013	25.09.2013	CTD	10	bdl	262*	WC
		MUC				All
		GC				GC-All
November 2013	08.11.2013	CTD	12	163	13	WC
		MUC				All
March 2014	13.03.2014	CTD	4	209	41*	WC
		MUC				All
June 2014	08.06.2014	CTD	7	47	61	WC
		MUC				All
September 2014	17.09.2014	CTD	13	bdl	234	WC
		MUC				All

1092 oxygen (O_2) and dissolved methane (CH_4) concentration

MUC = multicorer, GC= gravity corer, CTD = CTD/Rosette, bdl= below detection limit (5μM), All = methane gas
 analysis, porewater analysis, sediment geochemistry, net methanogenesis analysis, hydrogenotrophic
 methanogenesis analysis, GC-All= analysis for gravity cores including methane gas analysis, porewater analysis

methanogenesis analysis, GC-All= analysis for gravity cores including methane gas analysis, porewater analysis,
 sediment geochemistry, hydrogenotrophic methanogenesis analysis, WC= Water column analyses including
 methane analysis, chlorophyll analysis

1098 **Concentrations from the regular monthly Boknis Eck sampling cruises on 24.09.13 and 05.03. 14 (www.bokniseck.de)

1108 Figure 1



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0.14 Net MG Methanogenesis [mmol m⁻² d⁻¹] 0.10 0.00 m⁻² d⁻¹] 80.0 0.04 0.07 0.05 Hydrogenotrophic MG 0.00 Sept Sept Nov Mar Jun Mar Jun '13 '13 '13 '13 '14 '14 '14

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Figure 8









