Microbial communities from Arctic marine sediments respond slowly to methane addition during ex situ incubations

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Summary
Anaerobic methanotrophic archaea consume methane in marine sediments, limiting its release to the water column. However, their responses to changes in methane and sulfate remain poorly constrained. To address how methane exposure may affect microbial communities and methane- and sulfur-cycling gene abundances in Arctic marine sediments, we collected sediments from offshore Svalbard that represent three geochemical horizons where anaerobic methanotrophy is expected to be active, previously active, and long-inactive. Sediment slurries were incubated at in situ temperature and pressure with different added methane concentrations, and then studied for extended incubation times. Rapid increases in sulfur-cycling Deltaproteobacteria occurred after 30 days in previously active and long-inactive sediments. However, these sediments showed no evidence of methanotrophy after nearly eight months of incubation. Sediments from an active area of seepage began to reduce sulfate in a methane-dependent manner within months, preceding increased relative abundances of anaerobic methanotrophs ANME-1 within communities. However, methane did not structure microbial community changes across any incubation time or sediment type. These results suggest that active anaerobic methanotrophic populations may require years to develop, and that microbial community composition may affect methanotrophic responses to potential large-scale seafloor methane releases in ways that may provide insight for future modeling studies.

Originality-significance statement
Microbial communities that consist of anaerobic methanotrophic archaea, sulfate-reducing bacteria, and many others remove most of the methane produced in marine sediments worldwide, but are thought to develop over timescales of years and are challenging to study in situ. In this novel study, we describe concomitant observations of microbial community composition, methanotrophic activity, and abundances of methane- and sulfur-cycling genes to characterize responses of distinct sediment zones to varying methane concentrations across timescales of months. Our finding that microbial community changes precede growth and activity of methanotrophic populations illuminates the role of microbial community dynamics on methane flux to the overlying hydrosphere.

Introduction
Globally, marine sediments are sources and sinks of tens to hundreds of teragrams of methane per year (Valentine, 2002). Gas hydrates, vast reservoirs of temperature-sensitive methane, can be found in Arctic sediments below only hundreds of meters of water depth, and thus are expected to be influenced more quickly by warming than hydrates at lower latitudes (Hunter et al., 2013). Offshore western Svalbard, models of hydrate stability based on ocean warming trends predict increases in seafloor methane flux from slope and shelf regions (James et al., 2016 and references therein). Seafloor methane leakage along Arctic continental margins (Shakhova et al., 2010; Mau et al., 2017), has triggered concerns that warming bottom waters will destabilize hydrates, increasing methane influx into the water column or even the atmosphere (Westbrook et al., 2009). Though there is no evidence that this is presently occurring (Berndt et al., 2014; Hong et al., 2017; Wallmann et al., 2018), methane production,
consumption, and transport in Arctic seabed environments is complex and spatiotemporally variable.

Anaerobic methane oxidation (AOM) is a crucial biofilter that prevents up to 90% of this methane from reaching the hydrosphere (Reeburgh, 2007; Boetius and Wenzhöfer, 2013). This process, mediated by several clades of anaerobic methanotrophic archaea (ANME) and distributed in marine sediments worldwide, uses sulfate as an electron acceptor:

$$\text{CH}_4 + \text{SO}_4^{2-} \rightarrow \text{HCO}_3^- + \text{HS}^- + \text{H}_2\text{O} \quad \text{(Equation 1)}$$

To accomplish AOM, ANME often associate with sulfate-reducing bacteria (SRB) at sulfate-methane transition zones (SMTs), sediment horizons where sulfate reduction (SR) and AOM remove sulfate and methane from porewaters (Knittel and Boetius, 2009). Their global ubiquity and distribution within methane-rich sediments (Nunoura et al., 2008; Ruff et al., 2015) shapes microbial community structures in these zones (Harrison et al., 2009), and in situ observations suggest that these methanotrophic communities develop on timescales of years (Ruff et al., 2018; Klasek et al., 2019). Processes that could alter the distribution of methane and sulfate throughout the sediment column, such as changes in subseafloor methane flux (Hong et al., 2016), emission of mud breccia flows through mud volcanism (Ruff et al., 2018; Klasek et al., 2019), and sediment gravity flows (Hensen et al., 2003), all presumably impact microbial community structure, ANME/SRB populations, and ultimately AOM rates.

How ANME and other sediment microbial community members respond to changes in methane fluxes remains unclear, and represents a critical knowledge gap for understanding how a large-scale seafloor methane release could be mitigated in high-latitude regions (Dale et al., 2008). Though ANME have not been isolated in pure culture (Wegener et al., 2016), enrichment studies have characterized their doubling times on the order of months (Nauhaus et al., 2007). Incubations at elevated pressure, which increases methane solubility in the aqueous phase, have successfully stimulated AOM in a methane-dependent manner (Nauhaus et al., 2002; Deusner et al., 2010), enriched microbial biomass (Girguis et al., 2005; Wang et al., 2014), and characterized differences in temperature and pressure optima between ANME subpopulations (Nauhaus et al., 2005; Timmers et al., 2015; Bhattarai, Zhang, et al., 2018).

Our objective was to determine how microbial communities and ANME and SRB numbers would respond to changing methane fluxes in Arctic marine sediments in a controlled incubation. We collected marine sediments from seafloor gas hydrate mounds (GHMs) at Storfjordrenna, offshore Svalbard (Fig. 1) that emit methane into the overlying water and lie at depths corresponding to upper limits of gas hydrate stability (Serov et al., 2017). Three sediments with varying inferred AOM activity (Hong et al., 2017) were amended with methane and incubated at in situ temperature and pressure for up to eight months. We anticipated that 1) highest community changes would be observed in PA and IA sediments as microbial communities adjusted to geochemical perturbations; 2) SR-AOM rates would depend on added methane concentrations and take longer to establish in PA and IA sediments; and 3) correlations would be seen between SR-AOM rates and ANME/SRB marker gene abundances.

**Results**
To investigate how methane concentrations structure different microbial communities and control SR-AOM rates, we conducted several long-term (several month) microcosm incubations of Arctic marine sediments that varied with respect to their predicted AOM capability. We first consider in situ geochemical distinctions between each of the sediment types, and then discuss geochemical changes observed during incubations. These provide context for subsequent assessment of microbial community and gene abundance changes during incubation.

Incubated sediments were selected to reflect predicted patterns of spatiotemporal variability in AOM activity across the Storfjordrenna GHM area, based on rate data derived from transport-reaction models inferred from shapes of porewater sulfate profiles (Borowski et al., 1996; Dale et al., 2008). At Storfjordrenna, increases in subsurface methane flux that coincide with seepage atop several GHMs have been attributed to venting of deep reservoirs (Hong et al., 2017). A push core sample from an active site of seepage (PC1029 at 30-40 cm, from GHM3) was collected from just below a sediment depth where remaining porewater sulfate indicated incomplete reduction and directly atop gas hydrate nodules recovered at 45 cmbsf (Fig. 1B).

Methane phase transitions and an abrupt decrease in porewater sulfate with depth throughout this core suggest this horizon was undergoing AOM at the time of sampling, and we designate this sample as active (“A”) for future reference. At the southern flank of GHM3, a sample from 80-90 cm at GC1045 lies just below a sulfate-methane transition where alkalinity has increased through the sulfate reduction zone in an equimolar fashion (Fig. 1C). The curvature of the sulfate profile is consistent with a recent increase in methane flux at this site. A modeling study of a core from the same area (Hong et al., 2017) detailed an increase in methane flux pushing the SMT upwards at a rate of approximately 10 cm per year. This suggests the GC1045 80-90 cm sample may have been mediating AOM only a few years before sampling, and we designate this sample as previously active (“PA”). At GC1036, away from the GHMs, sulfide was not detected and porewater alkalinity was elevated only slightly above seawater values (Fig. 1D).

The gradual decrease in porewater sulfate concentrations points towards a low level of methane flux at this location. The redox chemistry at 20-35 cm appears to be uninfluenced by any recent methane incursion, suggesting this zone has not recently undergone AOM. We refer to this sample as inactive (“IA”).

When A sediments were incubated at in situ pressure (experimental setup shown in Fig. S1), a methane-dependent increase in media sulfide and a concomitant decrease in sulfate were observed after 118 days, with even stronger changes shown in the later stages of the 222-day incubation (Figs. S2 & S3). Incubations amended with methane showed similar increases in media DIC (Fig. S4). These observations indicate that porewater chemistry changes due to stimulation of AOM occurred sometime between two and four months after incubations began. Methane-free controls allow us to quantify baseline sulfate reduction rates attributed to organic matter oxidation, as others have done (Nauhaus et al., 2005; Deusner et al., 2010; Zhang et al., 2010). Although some SR was coupled to oxidative breakdown of organic matter, it never exceeded 62% of the rate measured in corresponding incubations with 5 mM CH₄ (A sediments from 0-101 days, Fig. 2). These data indicate that a significant fraction of the sulfate is consumed by AOM. SR rates for intervals of 118- and 222-day incubations increased with time and added methane, and exceeded rates measured in other sediment incubations (Fig. 2).
Stoichiometrically, sulfate consumed an average of 98% of the added methane in the A incubations, far more than in the other sediments, but the $\text{SO}_4^{2-}/\text{CH}_4$ ratio ranged from zero (in incubation stages where sulfate did not decrease) to 2.3 during the 30-101 days incubation interval of sample A amended with 1.5 mM CH$_4$. SR rates exceeded sulfide production rates in incubations with added methane, indicating precipitation of sulfide minerals. During AOM-SR, the $\delta^{13}$C-DIC pool becomes more depleted due to preferential selection of $^{12}$C-CH$_4$ by methanotrophs (Borowski et al., 1997), with a characteristic kinetic isotope fractionation effect ($\varepsilon$). We estimated a value for $\varepsilon$, and show that during our incubations of A samples it decreases from 42‰ to 19‰ in a methane-dependent manner (Fig. 3). Values of $\varepsilon$ below 30 are generally indicative of AOM (Whiticar, 1999).

In contrast to the A incubations, the IA and PA sediment microcosms showed little to no sulfide buildup or sulfate consumption (Figs. S2 & S3, Table 1) and there is no evidence of methane-dependent SR at any point during the timeseries (Fig. 2). Values of $\varepsilon$ did not reliably decrease when amended with 5 mM methane (Fig. 3, Table 1). Collectively these data show that AOM was not detected, even after nearly eight months, in IA and PA incubations.

Microbial communities from the three sediment types differed before incubations began (Table S1A). The A sediment communities consisted of mainly Epsilonproteobacteria and ANME (within class Methanomicrobia), while the IA was dominated by Deltaproteobacteria and the PA by Atribacteria (Fig. 4). PA sediments, sampled from below a sulfate reduction zone, were comparatively lower in OTU richness than IA (Fig. S5), in line with patterns of community diversity and assembly documented across SMTs (Starnawski et al., 2017). ANMEs constituted 5% of the PA sediment community, and were not detected in the IA sediment. Nevertheless, many overlapping classes were common to all three communities. Highest richness was seen in the IA sediment community (Fig. S5). During sample storage at 4°C after collection, IA and PA sediment communities showed reduced alpha diversity and increases in percent abundances of Deltaproteobacteria, but the A community did not change appreciably over nearly a year (Fig. S5).

Surprisingly, addition of methane did not broadly alter community structure in any sediment type (all p>0.6, Table S1B). Methane addition significantly ($\alpha=0.05$) increased percent abundances of only three OTUs that comprised over 1% of their communities. These included Draconibacterium and Desulfhorpalus in A sediment communities, and a Desulfuromonas in the PA that was most similar to D. svalbardensis, a psychrophile that couples acetate oxidation to Fe (III) reduction (Vandieken, 2006).

Sediment microbial communities, particularly the IA and PA, shifted noticeably after only 30 days of incubation, regardless of added methane concentration (Fig. 5). These changes first included increased percent abundances of Deltaproteobacteria, while eventually Clostridia and Bacteroidia became more dominant in longer incubations (Fig. 4). Longer incubations saw continued community changes in all but one instance (PA sediments between 118 and 222 days, Fig. 5). In contrast to the patterns of SR-AOM observed during incubation, the IA sediment communities changed the most over time, and the A communities the least (Table 1). Though IA
and PA communities incubated for 222 days appear on the right side of the NMDS plot (Fig. 5),
you still show discrete clustering by sediment type. Thus, across all incubations, we observe
little evidence of convergence towards a common community structure.
Across all sediment types, percent abundances of twenty highly abundant OTUs changed when
incubated for different times (Fig. 6). Many of these dynamic OTUs belong to the class
Deltaproteobacteria and are implicated in sulfur cycling in anoxic marine sediments. Some
(unclassified Desulfuromonadales) increased or decreased depending on the sediment type,
while in other instances, different OTUs within the same genus (Desulfocapsa OTUs 4 and 5)
display different concurrent responses within the same community. ANME percent abundances
showed time-dependent changes, with ANME-1b increasing in the A sediment communities
between 30 and 118 days, and ANME-2a-2b decreasing in the PA communities from 118 to 222
days. Percent abundances of some taxa, particularly Deltaproteobacteria, increased as samples
were stored at 4°C and atmospheric pressure prior to incubation (Fig. S5), and these largely
concur with the community shifts seen in the first month of incubations (Fig. 4). Two genera
from the PA communities, Desulfuromonas and Lutibacter, increased significantly during
storage. This made it appear they had decreased over the final several months of
incubations, so OTUs belonging to these groups were removed from Fig. 6.

The genes mcrA and dsrAB were quantified to represent abundances of anaerobic
methanotrophs (Luton et al., 2002; Hallam et al., 2003) and dissimilatory sulfate reducers
(Leloup et al., 2004), respectively. Before incubation, mcrA concentrations across the three
sediment types differed by several orders of magnitude, with the IA sediments showing the
lowest, and the A sediments showing the highest levels of this functional gene (Fig. 7A).
Differences in dsrAB genes were less notable, but lowest concentrations were seen in the PA
sediments (Fig. S6A). Together, these patterns in gene abundances are consistent with the
commonly understood spatial/redox distribution of methane- and sulfur-cycling microbes
through the sediment column.

During incubation, mcrA gene abundances only increased with incubation time and methane
concentration in PA sediments. Because ANME did not increase among PA incubations, this
could be attributed to slight increases in canonical methanogenic taxa (non-ANME
Methanomicrobia) from 0.05 to 0.2% of PA communities over the incubation period (p=0.0254,
one-way ANOVA). Methanococcoides was the most abundant methanogenic genus throughout
PA communities (0.06% on average), but neither it nor other genera showed clear time- or
methane-dependent changes in percent abundance. mcrA concentrations in IA sediments did
not change over incubation time (Fig. 7A, Table 1) and did not correlate with percent
abundances of ANME or canonical methanogens, the latter of which averaged only 0.05% of IA
communities. Changes in dsrAB abundances were more dynamic: IA incubations showed time-
dependent increases, while the PA incubations showed increases over the first 30 days followed
by a slight decline at 222 days (Fig. S6A). Though A communities did not show significant
changes in mcrA or dsrAB numbers throughout all incubation times and methane
concentrations, percent abundances of ANME-1 OTUs increased in a methane-dependent
manner after 222 days, while those belonging to ANME-2 showed little change (Fig. 7B).
A sediment communities, ANME comprised 13% of sequences, 80 times more abundant than canonical methanogens.

In contrast to A incubations, IA and PA communities showed negligible changes in ANME (<0.1% abundance) when amended with methane. ANME-3 were either of insignificant abundance or not detected across the entire dataset. A positive correlation was observed between SR rates and mcrA gene abundances in the A samples incubated for 118 or 222 days (Fig. 7C). Interestingly, we found no correlation between SR rates and dsrAB gene abundances (Fig. S6B). Rather, the methane concentrations supplied to these incubations appeared to limit SR-AOM rates to a greater extent than the levels of either gene.

Discussion

Anaerobic methanotrophic archaea consume methane produced in marine sediments, limiting its release to the water column, yet their response to changes in methane and sulfate remain poorly constrained. To better understand how methane concentrations influence microbial community structure and activity under in situ conditions, we incubated three sediments of varying predicted AOM capability at in situ pressure and with amended methane concentrations on months-long timescales. We open the discussion by considering how changes in methane supply over time influence incubation geochemistry and activity as measured by SR-AOM rates, which provides subsequent context for how these variables alter microbial community dynamics. We finally discuss how this information derived from an ex situ laboratory incubation study can be applied to analyze AOM dynamics and microbial communities in seafloor environments.

Incubation geochemistry

As $^{12}$C-methane is preferentially oxidized to DIC by microbes in marine sediments, the residual methane pool becomes isotopically heavier, resulting in a kinetic isotope effect ($\varepsilon$) ranging from 4-30‰. In contrast, acetoclastic methanogenesis produces $\varepsilon$ values ranging from 35 to 55‰ (Whiticar, 1999). The A incubations that showed high SR rates and a decrease in $^{13}$C in the DIC pool suggest anaerobic methane oxidation, with $\varepsilon$ values ranging from around 37‰ to as low as 19‰ in the late-term incubations amended with 5 mM methane (Fig. 3). These depletions occurred despite resupplying each incubation interval with methane from a tank, which had a CH$_4$ composition of -35‰. In the PA incubations, the decrease in $\varepsilon$ seen after 60 days may signal a slight shift towards AOM, but without any observation of methane-dependent SR, organoclastic SR may be more likely. Other reports of incubated microbial consortia mediating SR-AOM resulted in depletion of $\delta^{13}$C-CH$_4$, which was attributed to co-occurring methanogenesis (Seifert et al., 2006) and/or carbon isotope equilibration during sulfate limitation (Yoshinaga et al., 2014). However, because we did not observe depletion of sulfate (below 1 mM) or relative increases in methanotrophic taxa, these two processes likely did not occur in our incubations.

Because methane solubility in seawater increases with pressure, prior incubations of methane-rich seafloor sediments have shown that AOM rates increase with methane partial pressures over a broad timescale, ranging from days to months (Nauhaus et al., 2002; Meulepas et al., 2009; Deusner et al., 2010; Zhang et al., 2010). In most of these studies, no lag times were
observed before AOM was stimulated, though Meulepas et al. (2010) suggested that sulfate reduction coupled to organic matter breakdown may delay the onset of AOM during enrichment (Meulepas et al., 2009). We see a similar lag in initial stages of 222-day incubations of A sediments, where sulfate reduction rates are comparable among all methane treatments, but in later stages of A incubations, sulfate reduction rates were higher with added methane (Fig. 2).

In contrast to what we observed in our PA sediment incubations, AOM has been stimulated in previously oxic, non-seep sediments amended with methane after 24 weeks of continuous-flow incubation (Girguis et al., 2003). Overall, enrichment studies have yielded maximum SR-AOM rates that vary several orders of magnitude, from 0.1 to 286 µmol g dry weight\(^{-1}\) day\(^{-1}\) (Meulepas et al., 2009). Assuming a sediment porosity of 0.5 and a density of 2.6 g/ml, our A incubation maximum measured rate of 0.91 µmol g dry weight\(^{-1}\) day\(^{-1}\) appears on the low end of this range, but is comparable to other long-term incubations where ANME were enriched (Bhattarai et al., 2018). SR-AOM rates from our incubations are comparable to, or even an order of magnitude higher than, the numerically-derived rates from Vestnesa Ridge, west of Svalbard (Hong et al., 2016) despite the fact that methane concentrations in our incubations remained far below saturation. This may reflect different flow regimes at these sites, particularly between the A samples collected from an active seep and the Vestnesa SMTs that have lower fluxes of sulfate and methane. Other discrepancies in incubation rate measurements may be attributed to different methods used in sediment storage and rate calculation (Krüger et al., 2008), variations in methane and/or sulfate supply between batch and flow-through incubations (Zhang et al., 2010), physical parameters including pressure and temperature (Nauhaus et al., 2002), and variability in microbial community composition.

Methane-associated microbial community changes

We observed a limited effect of methane on microbial community structure across timescales of months in incubations that simulated seafloor conditions. Total percent abundances of ANME-affiliated reads increased from 8% to 15-20% in the A communities amended with either 1.5 or 5 mM methane, but differences were not apparent until 222 days (Fig. 7B, Table S3). After several months of high-pressure incubations targeting AOM, similar studies reported increases in ANME 16S genes (Girguis et al., 2005), cells and aggregates (Zhang et al., 2014), and percent abundances in archaeal communities from <1% to up to 50-60% (Aoki et al., 2014; Bhattarai et al., 2018). Years of enrichment with methane and sulfate were required for ANME and SRB to attain near-dominance of archaeal and bacterial communities (Wegener et al., 2016). Earlier studies that enriched ANME noted doubling times varying from 1.1 to 7.5 months (James et al., 2016, and references therein). The A incubations that exhibited highest SR-AOM rates showed increases in mcrA; assuming one copy of mcrA per ANME genome (Haroon et al., 2013), these increases translate to doubling times of 7.6 months which are similar to previously observed doubling times of up to seven months (Nauhaus et al., 2007). Known methanogens, including recently established methane-cycling clades such as Bathyarchaeota (Evans et al., 2015) and Verstraeetearchaeota (Vanwonterghem et al., 2016), were nearly two orders of magnitude less abundant than ANME in A communities, supporting the validity of this calculation. In contrast to (Zhang et al., 2014), who saw bacterial alpha diversity increase after incubations, we observed no such diversity changes associated with incubation (data not
shown) and even saw a decrease in alpha diversity associated with storage time in A_t and B_t samples (Fig. S5).

In the A sediment communities, *Draconibacterium* and *Desulforhopalus* OTUs proliferated in response to methane addition. *Draconibacterium* is a recently-described genus of facultative anaerobes from marine environments capable of fermenting polysaccharides (Du *et al.*, 2014; Li *et al.*, 2016). Members of *Desulforhopalus* reduce sulfate with lactate, propionate, alcohols, or aminosulfinic acids (Isaksen and Teske, 1996; Lie *et al.*, 1999), and thus their presence may reflect organoclastic sulfate reduction concurrent with SR-AOM.

**Temporal community changes**

Microbial community structure in all sediment types changed with the length of time they were incubated (Fig. 5) and not with the amount of methane they were amended with (Table S1). Nevertheless, shifts in microbial community structure were highest in samples from above or below SMTZs, which may reflect potential geochemical redox changes associated with incubation: *IA* sediments were introduced to sulfide in seawater media, while *PA* sediments were re-exposed to sulfate. Proliferation of an ANME-1b OTU between 30 and 118 days in T sediments (Fig. 6) coincides with, and may account for, the increase in AOM-SR rate during this interval (Fig. 2). Sulfide never accumulated over 3 mM in media from *IA* or *PA* incubations (Fig. S2), so microbial communities likely did not experience sulfide toxicity, which has been noted to occur around 5 mM or more (Maillacheruvu and Parkin, 1996). Increases in dsrAB among *IA* and *PA* 30-day incubations (Fig. S6) probably reflect the proliferation of sulfate-reducing clades of Deltaproteobacteria (Fig. 4), but they apparently mediate different cycles or do not reduce sulfate until later stages of incubation. Alpha- and Gammaproteobacteria, which possess oxidative types of dsrAB (Müller *et al.*, 2015), could also partially account for the increase in dsrAB. Because no sulfate reduction was observed at 30 days, high abundances of sulfur-cycling taxa may reflect other processes such as sulfide oxidation with organic matter (Heitmann and Blodau, 2006). Though mcrA abundances increased most quickly in *PA* incubations amended with highest amounts of methane (Fig. 7A), ANME percent abundances did not change appreciably (Table 1). Instead, this increase could reflect growth of methanogens, which can be active even in methane concentrations of several mM (Lazar *et al.*, 2012). OTUs belonging to the methanogenic clade *Methanococcoides* increased very slightly, from 0.04% to 0.21% after 222 days. *Methanococcoides* also increased up to 0.4% in *IA* communities, where ANMEs were only sporadically detected. Though some of the mcrA detected could reflect non-Euryarchaeotal lineages (Evans *et al.*, 2015; Vanwonterghem *et al.*, 2016), these recently-described potential methane-cycling phyla were very sparsely detected (Bathyarchaeota) or nonexistent (Verstraetearchaeota) in our communities.

The *IA* communities saw growth of OTUs belonging to *Geopsychrobacter* and *Desulfuromonas* (Fig. 6). These genera are capable of using acetate and a variety of other organic compounds derived from fermenters as electron donors, and Fe (III), Mn (IV), or S0 as electron acceptors (Pfennig and Biebl, 1976; Roden and Lovley, 1993; Holmes *et al.*, 2004). Since Fe (II) concentration peaked at over 19 µM within the *IA* sediment depth, this Fe (II) produced
through iron reduction could precipitate sulfide as iron sulfide minerals and account for the lack of sulfide accumulation observed in corresponding incubations (Fig. S2).

The A communities changed over time, but to a lesser degree than reported in a five-year enrichment with methane at ambient pressure (Aoki et al., 2014). No changes in mcrA or dsrAB abundance were observed in these incubations. This suggests that communities were already acclimated for AOM, and also that approximately eight months of incubation did not stimulate much ANME/SRB growth at the methane concentrations and conditions provided. However, a positive relationship between SR-AOM rate and mcrA abundance appears dependent on methane addition (Fig. 7C). In two outliers, mcrA exceeded $10^8$ copies g$^{-1}$, suggesting growth of ANME or other methanotrophic taxa that contain this gene. However, in the other incubations AOM-SR instead appeared to be driven by an increase in ANME activity.

Community activity in situ

Several factors need to be considered when interpreting incubation studies to infer in situ microbial community function and dynamics. Dilutions and “bottle effects” (Hammes et al., 2010) can alter bacterial community activity (Stewart et al., 2012) possibly due to organic matter flocculation (Pernthaler and Amann, 2005). We observed an unexpected proliferation of Bacteroidia and Clostridia in our incubations, considering each class comprised around 0.01% of sequences from cores collected in situ at the same GHM area (data not shown). Methanococcoides, Geopsychrobacter and Draconibacterium were also barely detected in situ, which suggests that conditions pertaining to incubation, potentially related to organic carbon turnover, may influence community composition. Though we replaced media periodically to limit sulfide accumulation, continuous-flow high pressure enrichment strategies would further minimize buildup of toxic metabolites (Zhang et al., 2010). These approaches, combined with high-pressure sediment coring, better approximate in situ physical and geochemical conditions while allowing experimental manipulation (Case et al., 2017).

Our finding that microbial communities, when supplied with methane and sulfate, may change significantly over several months without any apparent AOM activity suggests that incubations aiming to understand details of AOM adaptation under different conditions would likely require timescales of years. Field studies combining community analysis with seafloor observatory data across recent eruptions at a seafloor mud volcano found that AOM communities develop on timescales of 2-5 years (Ruff et al., 2018). Across a permafrost thaw gradient associated with decade-scale changes in vegetation, shifts in methanogenic communities and the dominance of acetoclastic methanogenesis have been associated with increased atmospheric methane fluxes (McCalley et al., 2014). In contrast, active communities of aerobic methanotrophic bacteria in wetland soils are capable of changing composition within a week after exposure to higher methane concentrations (Knief et al., 2006). In our incubations, increased percent abundances of several clades of sulfate-reducing bacteria appear to represent a stage that precedes AOM establishment that is consistent with field observations. The lack of activity in PA sediments is surprising, considering that ANME are present and the sediment likely corresponded with a SMT only years before sampling (Hong et al., 2017). In contrast, sediments from an active seep showed increases in SR-AOM rates that corresponded with higher abundances of mcrA genes and percent abundances of ANME. Microbial community data may be of use when paired with
geochemical markers to further elucidate methane seepage in areas with spatiotemporally heterogeneous methane fluxes (Hong et al., 2018).

Model-based approaches concerning changes in the activity or efficiency of the subseafloor microbial methane filter, possibly from increasing subseafloor methane fluxes (Dale et al., 2008; Boetius and Wenzhöfer, 2013), should consider realistic timescales for microbial community responses within particular sediment horizons. Our interpretations that zones previously capable of AOM may remain dormant after months of incubation with methane and sulfate suggest longer-term characterization of these lag times is needed to fully understand the development of methanotrophic communities in sediments. We further posit that *ex situ* incubations allow cross-validation of complementary field studies on AOM development, and aid in determining causative relationships between microbial community structure and biogeochemical function.

Fundamentally, it can be quite challenging to conduct controlled long-term experiments in the field, and these experiments may also be subject to some bias. By using laboratory experimentation to study microbial communities under defined conditions, such as increases in temperature (Fuchs et al., 2016) or organic matter concentrations (Babbin et al., 2016), novel insight can be gained into causal relationships among biogeochemical processes that exist in nature, and how these processes might be expected to change in response to environmental stimuli. Ideally, both controlled laboratory incubations and careful experiments performed at field sites can function together as complementary approaches to resolve difficult microbial ecology issues.

**Experimental Procedures**

**Fieldwork and sample collection**

Sediment samples were collected aboard the RV Helmer Hanssen on cruise CAGE16-5, from June 16th to July 4th, 2016, at the mouth of the Storfjordrenna trough fan, offshore Svalbard (Fig. 1A) with gravity cores or push cores. Gravity core (GC) 1045 was recovered from the south slope of GHM3, while GC1036 was collected several km east of GHM3 in 394 m water depth. Once cores were recovered, their plastic liners were removed from the core barrel, sectioned into 1 m segments, labeled, and split in half with a table saw to obtain working and archive halves. Core halves were stored horizontally at 4°C for up to two hours while alkalinity measurements were taken. Replicate PVC push cores (PC1029) for geochemical and microbiological sampling were collected approximately 30 cm from observed bubble seepage at GHM3 using a Sperre Subfighter 30k remotely operated vehicle (ROV) from the Centre for Autonomous Marine Operations and Systems (AMOS) equipped with a video camera and a raptor arm. Cores were extruded and subsampled using PVC tubing.

Sample selection for microbiology targeted discrete sediment horizons where AOM was predicted to be inactive (IA), previously active (PA), and active (A). These designations were inferred through porewater sulfate profiles (Fig. 1B-D), wherein nonlinear decreases in sulfate with depth have been attributed to the encroachment of AOM into shallow sediment horizons as a result of increased subsurface flux (Hong et al., 2017). These three samples corresponded roughly to geochemical zones located above, below, and within SMTs, whose depths were
inferred by onboard porewater alkalinity titrations from the three cores and later bolstered by porewater sulfate profiles (Fig. 1B-D). The low-alkalinity inactive (IA) sample was collected from GC1036 at 20-35 cm below seafloor, and the previously active (PA) sample was taken from GC1045 at 80-90 cm depth, just below the porewater alkalinity increase corresponding to the SMT. The active (A) sample, at 30-40 cm from PC1029, was chosen because it lay below an alkalinity increase but above gas hydrates recovered at 45 cm (Fig. 1B).

Using ethanol-sanitized spatulas to scrape away the outer few mm of sediment from the working core half, sediment samples were placed into Gaspak anaerobic pouches with oxygen-scavenging catalysts (BD Biosciences) and stored in a nitrogen-filled bag at 4˚C until the cruise ended, then transported to Oregon State University on ice. Unincubated subsamples were frozen immediately after collection. Samples were stored in anaerobic pouches in a cold room for 100 to 242 days until incubations began. Anaerobic methanotrophic communities are thought to be resilient to pressure changes of these magnitudes (0.1 to 4 MPa); pressures up to 20 MPa have been required to significantly alter composition of enriched methanotrophic communities (Bhattarai, Zhang, et al., 2018).

**Incubation**

A manual pressure pump generator (High Pressure Equipment Co. part 87-6-5) was used to hydrostatically pressurize standard or customized pressure vessels to 4.0 MPa, approximating in situ pressure (Fig. S1A). At this pressure, and at an incubation medium salinity of 35 ppt, hydrate can form between 3.0 and 3.1˚C, provided sufficient methane concentration (Sun and Duan, 2007). In an anaerobic glove bag, sediment samples were homogenized, and 2.0 g sediment and 8.0 ml anoxic artificial seawater medium were added to cutoff Hungate tubes (Fig. S1B), similar to (Bowles et al., 2011). The medium, modified from (Widdel and Bak, 1992), contained 2 mM HCO₃⁻, 1.5 mM HS⁻, and 10 mM SO₄²⁻. Methane gas (99.7% pure) was sampled into a gastight ALTEF polypropylene bag (Restek) and transferred to glass cutoff Hungate incubation tubes (Bellco) using gastight syringes. Once fully dissolved, medium methane concentrations were 0, 1.5, or 5 mM for the different treatments. Tubes were placed upside-down in vessels, pressurized to allow methane to dissolve completely and instantaneously, and incubated at 4˚C for 30, 118, or 222 days. Incubation microcosms were conducted in triplicate.

222-day incubations were started in October 2016, while 30- and 118-day incubations began in February 2017. The cold room temperature fluctuated between 4-6˚C during storage and incubation, with a single excursion to 8˚C for several days in January 2017. During each week of incubation, pressure vessels were gently shaken to resuspend sediment slurries and repressurized to 4.0 MPa as necessary.

Medium from 118-day incubations was sampled once (at 61 days), and medium for 222-day incubations was sampled twice (at 101 and 188 days). After letting sediment settle, as much of the remaining methane gas and medium as possible was removed (4.5 to 7 ml) before replenishing with fresh methane and medium. Medium was sampled from incubation tubes with a needle and syringe and passed through a 0.2 µm filter. Samples for sulfate measurements were frozen at -20˚C in Eppendorf tubes. Medium for sulfide measurements was preserved using a saturated zinc acetate solution, centrifuged, and stored at 4˚C. Medium for dissolved inorganic carbon measurements was poisoned with a saturated mercuric chloride
solution. Sediment slurries were transferred to 15 ml falcon tubes, centrifuged, remaining supernatant poured off, and pellets frozen at -80°C.

**Geochemistry**

Sediment porewater was collected using rhizon samplers spaced 10-25 cm through the sediment column (or every 1-2 cm for PC1029). Total porewater alkalinity (TA) was titrated onboard. Depending on the expected TA, we used 0.1 to 0.5 ml of porewater for titration in an open beaker with constant stirring. pH was manually recorded with every addition of 1.2 mM HCl. Seven to ten measurements were performed for every sample. TA was calculated from the recorded pH, and the amount of acid was added using the Gran function. Details of the calculation were reported previously in (Latour et al., 2018). Increases in porewater alkalinity determined by onboard titrations were used to roughly constrain the SMT depths (within 30 cm) for sampling purposes. Total sulfide ($\Sigma$HS) concentrations from porewaters were measured spectrophotometrically using the method of (Cline, 1969). Samples were preserved onboard with a 23.8 mM zinc acetate solution onboard less than 30 minutes after the syringes were disconnected from rhizons, and then kept frozen until shore-based analysis. Dissolved iron (II) was determined spectrophotometrically at 595 nm onboard with a ferrospectral complex in 1% ascorbic acid. Details of sulfide and iron (II) analyses are also available in (Latour et al., 2018).

Porewater sulfate was analyzed with a Dionex ICS1100 ion chromatograph (IC) at the Geological Survey of Norway (NGU). An IonPac AS23 column was equipped on the IC with the eluent (4.5 mM NaCO₃ and 0.8 mM NaHCO₃) flow set to be 1 mL/min. Sulfide from incubation medium was analyzed by the USEPA methylene blue method (Association et al., 1989) using commercial reagents (Hach) and a Hach 4000 UV/Vis spectrophotometer set to 690 nm after calibration with a sulfide standard (Sigma). Sulfate from incubation medium was measured on an Integrion HPIC RFIC Ion Chromatograph (Thermo Fisher Scientific) using a Dionex IonPac AS18 Analytical Column 4X250mm column. A KOH buffer was used as an eluent, and ranged from 22 to 40 mM throughout the 20-minute run. Sulfate eluted at 10.5 min. Calibrations were prepared with five laboratory standards (Dionex), and replicate IC measurements varied on average by 0.6%. To obtain concentrations and $\delta^{13}$C isotopic signatures, DIC samples were added to Exetainer vials (Labco), purged with helium, mixed with orthophosphoric acid to release CO₂, equilibrated for 10 h, and measured using a GasBench-DeltaV system as explained in (Torres et al., 2005). $\delta^{13}$C values are expressed in ‰ relative to PDB. Sulfate reduction, sulfide production, and DIC production rates were calculated by taking the difference of the medium concentrations of these constituents across incubation intervals and dividing by the mass of bulk sediment and time interval incubated. For each interval, the amounts of sulfate reduced and methane added were compared stoichiometrically to determine the percentage of methane that may be oxidized by sulfate. $\delta^{13}$C-CH₄ was measured using a MAT 253 isotope ratio mass spectrometer outfitted with a ConFlo IV interface (all components Thermo Fisher Scientific Inc.) at MARUM, University of Bremen, Germany and determined to be -35‰ VPDB (T. Pape, personal communication). A $\delta^{13}$C-CH₄ of -35‰ was used for all $\varepsilon$ calculations because methane of the same isotopic signature was resupplied during incubation stages. Kinetic isotope effects (notated here as $\varepsilon$, epsilon) were determined by subtracting $\delta^{13}$C-CH₄ from $\delta^{13}$C-DIC measured after separate incubation intervals, and then corrected for the temperature-dependent carbon isotope fractionation between DIC and CO₂ at
4°C (Zeebe and Wolf-Gladrow, 2001) to allow comparison with previously determined $\varepsilon$ values (Whiticar, 1999):

$$\varepsilon = (\delta^{13}C{-\text{DIC}}) - (\delta^{13}C{-\text{CH}_4}) + 7 \quad \text{(Equation 2)}$$

### DNA Extraction, Amplification, Sequencing, and Analysis

DNA was extracted from sediments in a clean laminar flow hood using a Qiagen DNeasy PowerSoil kit (Hilden, Germany) following the manufacturer’s protocol. The Earth Microbiome Project 16S Illumina protocol was used to prepare amplicons for sequencing. Briefly, V4 regions of bacterial and archaeal 16S rRNA genes were amplified in triplicate 25 ul reactions using universal 515-forward and 806-reverse primers (Caporaso et al., 2011) modified with dual-indexed Illumina sequencing adapters (Kozich et al., 2013). The thermal cycling protocol of Caporaso et al. 2011 (Caporaso et al., 2011) was followed without modifications. After confirming amplification with agarose gel electrophoresis, triplicate PCR products were pooled and purified with a Qiagen QIAquick PCR purification kit. Amplicon concentrations were quantified with a Qubit fluorometer using the Qubit dsDNA high sensitivity assay kit and pooled in equimolar amounts. Illumina MiSeq V2 paired-end 250 bp sequencing was performed by technicians at Oregon State University’s Center for Genome Research and Biocomputing (CGRB). A sediment-free DNA extraction blank was amplified and included in the sequencing run. Raw fastq sequences were uploaded to the NCBI Sequence Read Archive (SRA) under the BioProject number PRJNA532941.

16S gene sequences were processed with mothur (Schloss et al., 2009) (v.1.39.3) following an established pipeline (Kozich et al., 2013). Reads were clustered into operational taxonomic units (OTUs) at a 97% similarity level and taxonomically classified using the SILVA database (v. 128) (Quast et al., 2013). We manually examined sequences known to be contaminants in DNA extraction kits and in subsurface ecosystems (Salter et al., 2014; Sheik et al., 2018). Comamonas represented the only suspicious genus, and because it was only present in an extraction blank and at low abundance, no contaminant sequences were removed from the dataset. After removal of singleton OTUs, communities were rarefied to 4,738 reads and relative abundances were calculated. Alpha diversity metrics (number of OTUs and Chao1, Shannon, and Simpson indices) were then determined. To compare beta diversity, a tree file containing the most common individual sequence from each OTU was constructed using clearcut (Evans et al., 2006). Weighted Unifrac (Lozupone et al., 2007) distances were calculated from the untransformed OTU table to obtain coordinates for non-metric multidimensional scaling (NMDS) ordination. ANOSIM (Clarke, 1993) was used to test for differences in community structure between incubation treatments relating to time or methane addition. Metastats (White et al., 2009) was used to determine whether individual OTUs were differentially abundant between communities corresponding to these treatments. Because microbial community composition can change appreciably during storage at these temperatures and on these timescales (Mills et al., 2012), unincubated sediments were subsampled and frozen at several times during storage to determine whether changes were brought about by incubation conditions specifically (Fig. S5). OTUs belonging to genera that changed by more than 1% in abundance during sediment sample storage were omitted from results that reported changes in OTU percent abundances.

### Droplet Digital PCR
Droplet digital PCR (ddPCR) was used to quantify abundances of functional genes *dsrAB* and *mcrA* using primer pairs described by Kondo et al. (Kondo et al., 2004) and Luton et al. (Luton et al., 2002), respectively. Reactions of 22 μl volume were prepared in a clean PCR hood in 96-well plates using 1x Bio-Rad QX200 ddPCR EvaGreen Supermix, 200 nM primers, and 0.88 μl of tenfold-diluted genomic DNA. Droplets were generated on a QX200 AutoDG Droplet Generator using automated droplet generation oil for EvaGreen Supermix (Bio-Rad). Thermal cycling was performed immediately afterwards on a Veriti 96-well thermal cycler. Protocols began with a single initialization step at 95°C for 5 min and then proceeded to 40 cycles of denaturation at 95°C for 30 sec, annealing for 1 min (at a temperature of 53°C for *mcrA* and 58°C for *dsrAB*), and for *mcrA* only, an extension at 72°C for 75 sec. Signal stabilization steps (4°C for 5 minutes, then 90°C for 5 minutes) were then performed before maintaining a 4°C hold. To ensure uniform heating of all droplets, the ramp rate for all amplification cycles was set to 2°C/minute. Reactions were held at 4°C overnight and read with the Bio-Rad QX200 Droplet Reader after 12 h. Droplet generation and reading were performed at OSU’s CGRB core facility. Normalization was performed by inspecting fluorescence distributions using QuantaSoft software (Bio-Rad). Threshold fluorescence values were manually imposed by visually inspecting distributions of DNA extraction blank and no-template-added control samples. Amplicon copy numbers per well were then converted to copies per gram bulk sediment.
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Table 1. Summary of key geochemical and microbiological measurements across all incubated sediment types that changed significantly based on added methane concentration [CH$_4$], incubation time, or both. ND: not detected. Because all methane-dependent changes observed were also time-dependent, methane concentration alone is not listed as a variable.

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<td>A</td>
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Figs. 2, S2-4  Fig. 3  Fig. 5  Fig. 7A  Fig. S6  Fig. 7B
Figure 1. Map of the Storfjordrenna trough mouth fan, south of Svalbard, showing locations of the three cores taken with white polygons indicating areas of methane seepage observed during sampling (A). Sulfate, sulfide, alkalinity, and iron (II) porewater profiles are shown successively in (B-D): PC1029, a push core from a seep at GHM 3; GC1045, a gravity core from the flank of the same GHM, and GC1036, a reference core near the mounds. Red bands within dotted lines indicate sediment sample depths collected for incubations. Sediment sample names correspond to inferred methanotrophic capacity: active (A), previously active (PA), and inactive (IA).
Figure 2. Sulfate reduction rates measured across incubation intervals. Panels are separated horizontally by total incubation time (118 or 222 days) and vertically by sediment type. Gray shades designate added methane concentrations, and error bars represent 95% confidence intervals from triplicate measurements. Methane-dependent SR was only observed in the A sediments collected from an area of active seepage.
Figure 3. Changes in $\delta$, the kinetic isotope fractionation during anaerobic oxidation of methane to DIC, over incubation time. Panels are separated horizontally by total incubation lengths (118 or 222 days) and vertically by sediment type, and gray shades designate methane concentrations. Decreasing values of $\delta$ during incubations of A sediment samples indicate a more prominent influence from AOM.
Figure 4. Percent abundances of class-level taxonomic divisions across all sediment types, times incubated, and initial methane concentrations. These sequences constitute >90% of the dataset. The extent of variation between duplicate or triplicate microbial communities incubated under the same conditions is conveyed by widths of black outlines.
Figure 5. NMDS ordination of weighted Unifrac distances between communities. Symbols are colored according to incubation length, and shaped according to sediment type. Polygons indicate statistically distinct community structures as determined by ANOSIM using $\alpha=0.05$ with a Bonferroni correction for multiple comparisons (Table S1). Initial methane concentrations are not indicated because methane did not structure community composition across any sediment type (Table S1).
Figure 6. Percent abundance and taxonomic identity of the most common OTUs (each >1% of their communities) whose abundances changed significantly (α=0.05) when incubated for different lengths of time. Blue dots represent incubations of different lengths, and are connected by black or red vertical lines that respectively show increases or decreases in OTU percent abundance over the corresponding interval. Where multiple OTUs share taxonomic identity, OTU numbers are included. Added methane concentrations are omitted from this figure, as they changed relative abundances of only three other OTUs (discussed earlier in text).
Figure 7. Log-normalized abundances of mcrA genes (g\(^{-1}\) bulk sediment) in incubated samples, enumerated by ddPCR (A). Points are shaded by initial methane concentrations, and X denotes readings below detection (10\(^3\) genes g\(^{-1}\)). One measurement was taken for preincubated (day 0) samples. Significant differences in gene abundances across incubation times and methane concentrations, excluding day 0 samples, are shown as brackets with corresponding p-values (two-way factorial ANOVA, \(\alpha = 0.05\)). Table S2 gives additional information on these tests. Orange box shows a subset of active (A) sediments incubated for 118 or 222 days. Of this subset, Panel (B) shows changes in percent abundances of ANME clades, with 95% confidence intervals shown for three replicates and brackets with asterisks denoting statistical significance among close comparisons. (C) relates sulfate reduction rates and mcrA gene abundances of the same subset through a linear regression.
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