Fine-scale community structure analysis of ANME in Nyegga sediments with high and low methane flux

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To obtain knowledge on how regional variations in methane seepage rates influence the stratification, abundance, and diversity of anaerobic methanotrophs (ANME), we analyzed the vertical microbial stratification in a gravity core from a methane micro-seeping area at Nyegga by using 454-pyrosequencing of 16S rRNA gene tagged amplicons and quantitative PCR. These data were compared with previously obtained data from the more active G11 pockmark, characterized by higher methane flux. A down core stratification and high relative abundance of ANME were observed in both cores, with transition from an ANME-2a/b dominated community in low-sulfide and low methane horizons to ANME-1 dominance in horizons near the sulfate-methane transition zone. The stratification was over a wider spatial region and at greater depth in the core with lower methane flux, and the total 16S rRNA copy numbers were two orders of magnitude lower than in the sediments at G11 pockmark. A fine-scale view into the ANME communities at each location was achieved through operational taxonomical units (OTU) clustering of ANME-affiliated sequences. The majority of ANME-1 sequences from both sampling sites clustered within one OTU, while ANME-2a/b sequences were represented in unique OTUs. We suggest that free-living ANME-1 is the most abundant taxon in Nyegga cold seeps, and also the main consumer of methane. The observation of specific ANME-2a/b OTUs at each location could reflect that organisms within this clade are adapted to different geochemical settings, perhaps due to differences in methane affinity. Given that the ANME-2a/b population could be sustained in less active seepage areas, this subgroup could be potential seed populations in newly developed methane-enriched environments.

Keywords: ANME, pyrosequencing, AOM, community structure, Nyegga, cold seep, stratification

INTRODUCTION

Anaerobic methanotrophs (ANME) play a vital role in the global carbon cycle budget, acting as methane sinks in marine systems. Through anaerobic oxidation of methane (AOM) they are estimated to consume >90% of the 85–300 Tg CH₄ annually produced, and thereby contribute to a strong reduction of methane emission to the atmosphere (Knittel and Boetius, 2009). Their main niche in marine sediments is the sulfate-methane transition zone (SMTZ) which is formed when methane from subsurface reservoirs meets sulfate penetrating from the water column through advection (Berelson et al., 2005; Treude et al., 2005b; Knittel and Boetius, 2009). The location of the SMTZ ranges from a few decimeters to several hundred meters below the seafloor, and is influenced by the local geological settings such as the depth of the methane production zone, the flux of methane, and sulfate through the sediment column and their consumption rates (Knittel and Boetius, 2009). Sequences from all the currently defined ANME clades, ANME-1, ANME-2, and ANME-3, have been retrieved from SMTZs from areas like the Gulf of Mexico (Lloyd et al., 2006), Skagerrak (Parkes et al., 2007), Haakon Mosby Mud Volcano (HMMV; Niemann et al., 2006) respectively, and the present understanding is that no specific ANME clade is considered characteristic for the SMTZ.

Although the SMTZ is often dominated by members of the ANME clade, other archaeal taxa besides ANME have been found to be enriched within the SMTZ. Uncultured Archaea like Marine Benthic Group B (MBG-B) [also named Deep Sea Archaeal Group (DSAG)], Miscellaneous Crenarchaeotic group (MCG), and Marine Benthic Group D (MBG-D) are among the most abundant taxa in systems like the Peru margin, Aarhus bay, Benguela Upwelling System, and Santa Barbara Basin (Sørensen and Teske, 2006; Schafer et al., 2007; Harrison et al., 2009; Webster et al., 2011). No ANME population was identified in sediments from the Peru margin and Cascadian margin in the study by Inagaki et al. (2006), despite the presence of gas hydrates located close to the SMTZ (present at 50–110 mbsf and below), while ANME was found among the low-abundance taxa in the other locations mentioned above. The highly abundant DSAG in Peru margin sediments has been suggested to be involved in the consumption of methane or in sulfate reduction (D’Hondt et al., 2004; Biddle et al., 2006; Inagaki et al., 2006), although the metabolic capability of this taxon remains unsolved. Furthermore, the methane...
fluid flux into the SMTZ in Peru margin has been estimated to 1.6–8.8 mmol/m² year (Biddle et al., 2006), which is considerably lower than the flux in ANME-dominated areas like Hydrate Ridge (11–33 × 10³ mmol/m² year; Torres et al., 2002), the Gulf of Mexico (500–2300 mmol/m² year; Lloyd et al., 2010), and Eckernförde Bay (240–690 mmol/m² year; Treude et al., 2005a). Hence, the microbial community composition and abundance of various archaeal taxa in marine sediments could be related to the supply of methane through the system over time, and whether the available methane is sufficient to sustain an ANME-dominated community over other uncultured Archaea.

For the G11 pockmark at Nyegga (the southern Voring Plateau, offshore Mid-Norway) located at water depth of 730 m the methane flux rate is estimated to 300–500 mmol/m² year (Chen et al., 2010). New high-throughput pyrosequencing technologies allow a deeper sampling of the ecosystems of interest, and approaches involving barcodes or unique DNA sequence identifiers have been developed for multiplex sequencing (Huber et al., 2007; Parameswaran et al., 2007; Hamady et al., 2008). The stratification of microorganisms in sediments from the G11 pockmark was recently analyzed using by FISH, quantitative PCR, and 16S rRNA gene amplicon libraries of several subsamples (Roalkvam et al., 2011). The sediment core was sampled inside the G11 pockmark in a pingo structure, which is characterized by an elevated seafloor due to the local accumulation of gas hydrates below (Howland and Svensen, 2006), where the methane flux is relatively high. The horizons in the shallower parts of the core [0–3 cm below seafloor (cmbsf)] were dominated by aerobic methanotrophs within Gammaproteobacteria, and sulfur oxidizing taxa within Epsilonproteobacteria. At depths below 4–5 cmbsf, a stratification of ANME clades was observed with transitions between horizons dominated by ANME-2a/b, ANME-1, and ANME-2c with increasing depth.

Here, we used 454-pyrosequencing of 16S rRNA tagged PCR-amplicons combined with quantitative PCR to investigate the stratification of the microbial communities in a Nyegga sediment with a relatively low methane flux 80 mmol/m² year and deep SMTZ at 205–255 cmbsf (Chen et al., 2011). Furthermore, we compare the microbial diversity and dominating ANME phylogenotypes in sediments with low methane flux with those present in sediments with higher methane flux from a pingo structure within the G11 pockmark (Roalkvam et al., 2011). Overall, we demonstrate that the local fluid flow regimes influence the abundance and diversity of microbial populations in cold seep sediments at Nyegga, and possibly in cold seep sediments in general.

MATERIALS AND METHODS

SITE DESCRIPTION AND SAMPLING

The Nyegga area is located on the upper Mid-Norwegian continental slope, at the northeast flank of the Storegga Slide (Figure 1), and is characterized by a high density of pockmarks and fluid seepage structures (Evans et al., 1996; Hustoft et al., 2010; Reiche et al., 2011). The area has been a field for multidisciplinary studies on gas hydrates, authigenic carbonates, fluid flow, and pore-water geochemistry the last decade with a special focus on the active micro-seeping area around pockmarks G11 and CN03 (also called CNE03; Howland et al., 2005; Howland and Svensen, 2006; Mazzini et al., 2006; Chen et al., 2010; Hustoft et al., 2010; Ivanov et al., 2010; Plaza-Faverola et al., 2010, 2011; Vaular et al., 2010; Reiche et al., 2011). Recent 2D/3D seismic and multibeam mapping of the Nyegga area has revealed an area with a high density of pockmark structures, many with underlying gas blanking areas extending down to a pronounced bottom simulating reflector (BSR) at 250–300 m depth below seafloor (mbsf; Bünz et al., 2003; Hustoft et al., 2007, 2009, 2010; Hjelstuen et al., 2010; Plaza-Faverola et al., 2010; Reiche et al., 2011). During a cruise with R/V G.O.Sars to the Nyegga area in August of 2008, the 3 m long gravity core GS08-155-15GC (referred to as 15GC hereafter) was retrieved from the CN03 area (64°45.274′N 05°04.088′E) at 725 m water depth. The ambient seawater temperature was measured with CTD to be between −0.6 and −0.7°C. After retrieval of 15GC, one half of the core was immediately sampled for detailed microbial diversity studies and geochemical measurements while the other half was stored at 4°C as an archive for non-destructive MST and XRF core scanner studies in laboratories on land. Rhizon samplers were used to extract pore-water from eight horizons throughout the core; at 24, 57, 89, 129, 171, 244, 258, and 290 cm below seafloor (cmbsf). The subsamples were preserved in glass vials and kept cool until they were analyzed according to the approach in Chen et al. (2010) in order to determine the concentration of dissolved sulfate (SO₄²⁻) and total dissolved hydrogen sulfide (ΣH₂S). Subsamples for DNA extraction were aseptically retrieved at 10, 30, 50, 80, 100, 120, 140, 160, 180, 200, 220, 240, 255, 270, and 300 cmbsf (±0.5 cm) by using sterile 1 mL tip cut plastic syringes before they were snap-frozen in liquid N₂ and stored at −80°C.

ESTIMATION OF METHANE FLUX

Sulfate gradients may be used to estimate the in situ methane flux (Borowski et al., 1996). The sulfate diffusive flux is obtained from the linear zone in the concentration profile, i.e., from the depth range in which there is no production or consumption. Fick’s first law (Krom and Berner, 1980) was used to calculate the sulfate diffusive flux in the core, as described by Chen et al. (2010). As the consumption of sulfate and methane has the stoichiometry 1:1 during AOM (Boetius et al., 2000), the sulfate diffusive flux is equivalent to the methane flux. The flux was estimated to ~80 mmol/m² year in 15GC, when a core porosity of 63% was used (Chen et al., 2011).

DNA EXTRACTION AND 16S rRNA GENE AMPICLON LIBRARY PREPARATION

Total genomic DNA was extracted from ~0.5 g of sediment from all subsamples using FastDNA Spin kit for soil (MP Biomedicals), and was subsequently quantified by A260/A280 ratio measurements, as described in Roalkvam et al. (2011). The pipeline of 16S rRNA gene amplicon library preparation, sequence filtering, and taxonomical classification of amplicons is described elsewhere (Lanzén et al., 2011; Roalkvam et al., 2011). In short, DNA from seven subsamples in 15GC (10, 30, 80, 120, 180, and 270 cmbsf) was applied in a two step PCR in order to generate a 16S rRNA gene amplicon library for each horizon where both PCRs followed a previously described protocol (Roalkvam et al., 2011). In order to evaluate the accuracy of the PCR and the sequencing step, subsample 270 cmbsf from 15GC was analyzed in triplicates.
Template DNA from all subsamples were amplified in triplicates using the primers Un787f (5′-ATTAGATACCCNGGTAG; Roesch et al., 2007) and Un1392r (5′-ACGGGCGGTGWGTRC; modified from Lane et al., 1985). The triplicates were pooled, and impurities were removed using MinElute® PCR purification kit (Qiagen). Purified amplicons were used as template in a second PCR where the above mentioned primers were modified to specifications in Lib-L chemistry: the GS FLX Titanium Primer A sequence and a specific MID sequence of 10 bp for each sample was included in forward primer Un787f, while the GS FLX Titanium Primer B sequence was included in reverse primer Un1392r. The final amplicons were purified as described above and the concentration was determined by SYBR-Green quantification, as described in Roalkvam et al. (2011). Prior to the pyrosequencing all samples were pooled, and a final purification using Agencourt AMPure XP (Beckman Coulter genomics) was applied to ensure removal of all impurities. The GS FLX instrument (Roche) at the Norwegian Sequencing Centre was used with 450 bp chemistry for 454-pyrosequencing of all amplicons. The raw sff-files of 16S tag-encoded amplicons from all subsamples in gravity core 15GC from Nyegga have been submitted to the Sequence Read Archive under the accession number SRA026733.

QUALITY FILTERING OF 16S AMPHILCON SEQUENCES AND TAXONOMIC CLASSIFICATION

Quality filtering and noise removal of pyrosequencing reads of amplicons were carried out using AmpliconNoise (Quince et al., 2011) as described in Roalkvam et al. (2011). In summary, noise, and errors introduced during PCR and pyrosequencing are corrected during four steps: filtering, flowgram clustering, sequence-clustering, and chimera removal. The filtered sequences were aligned to a reference database prepared from Silva SSURef release 102 (Lanzén et al., 2011) using blastn (default parameters). Sequences with a bit-score above 150 were assigned to their equivalent taxa in the modified Silva Taxonomy described above based on the taxonomy of the best blastn bitscores within a 10% range, using MEGAN version 3.7 (Huson et al., 2007). Finally, the assignments were exported and weighed according to its cluster’s copy number.

The amplicon library from 270 cmbsf was made in triplicates prior to sequencing to test the precision and reproducibility of our primers and the pipeline for amplicon construction and filtering. The relative abundance of the taxa listed at all taxonomical levels was compared between the parallel samples, all showing nearly the same relative taxa distribution (maximum deviation at any
given taxonomic level was 0.014%). The triplicates were therefore merged and treated as one subsample, comprising 61268 reads, due to the low deviation in relative abundance.

**OPERATIONAL TAXONOMIC UNIT ASSIGNMENT AND DIVERSITY INDEX ESTIMATIONS**

To compare the microbial communities in GC15 from the CN03 area with those in the more active area within the pingo structures at G11 pockmark (Roalkvam et al., 2011) on operational taxonomical units (OTU)-level, the individual 16S rRNA gene tagged amplicon files were merged and grouped into OTUs using the AmpliconNoise software and its incorporated features (Quince et al., 2011). In AmpliconNoise, a quick pre-clustering of flow-grams was performed prior to the pair-wise alignment of sequences using the Needleman–Wunsch algorithm (Needleman and Wunsch, 1970), followed by a hierarchical maximum linkage clustering with of 97% sequence identity. In order to assign a taxon to each cluster, one representative sequence from each OTU was selected and aligned to the Silva Taxonomy using MEGAN. The sequences representing each OTU were aligned to the Silva SSURef release 104 in Arb, and a total of 23 sequences were excluded due to short length (<220 bp) or chimeras, leaving 3299 sequences.

Diversity indices were calculated by the Shannon–Weaver index (Weaver and Shannon, 1949) and Rao’s quadratic entropy index (Rao, 1982) using R (version 2.13.1) with the Vegan package integrated or an in-house R script, respectively. The distance matrix needed for Rao’s quadratic entropy index was generated using Phylogeny Inference Package (PHYLIP; Felsenstein, 1989) in Arb (Version 5.0). Diversity indices, such as the Shannon index, may be influenced by sampling size. To compare diversity indices across pyrosequencing libraries of variable size, all sequence tags from all samples were first clustered into OTUs. Then each library were randomly subsampled using a subsampling size equal to the sample size of smallest library (796 reads) and by keeping the original OTU assignments for each sequence tag. Reported mean values and standard deviations were calculated from indices calculated from 1000 subsampling iterations per sample.

**QUANTITATIVE PCR**

The number of 16S rRNA genes from both Archaea and Bacteria in each subsample of 15GC were enumerated using real-time quantitative PCR, as described in Roalkvam et al. (2011). In short, genomic DNA from subsamples was quantified in duplicates, where each reaction (20 μl) contained 1 × Power SYBR-Green PCR Master Mix (Applied Biosystems), 1 μM of each primer, and 1 ng template. The 16S rRNA genes of bacterial origin were amplified using the primers B338f (5′-ACTCCTACGGGAGGCAGC; Ovreas et al., 1997) and A907r (5′-ATTACCGCGGCTGCTGG; Muyzer et al., 1993) and 40 cycles of the thermal program described by Einen et al. (2008). The standard curve was generated using DNA from Escherichia coli, and genomic DNA from Archaeoglobus fulgidus was used as negative control. Similarly, archaeal 16S rRNA genes were amplified using primers Un519f (5′-TTACCCTACGGGAGGCAGC; Amann et al., 1995) and B518r (5′-ATTACCGCGGCTGCTGG; Muyzer et al., 1993) and 40 cycles of the thermal program described by Roalkvam et al. (2011). The linearized fosmid 54d9 was used to generate standard curve and genomic DNA from E. coli was used as negative control. For each subsample, the mean number of 16S rRNA gene copies/g sediment and corresponding standard deviation was used as enumeration of Bacteria or Archaea in the core.

**RESULTS**

**GEOCHEMISTRY**

The concentration of dissolved sulfate (SO$_4^{2-}$) and total dissolved hydrogen sulfide (ΣH$_2$S) in pore-water from core 15GC from the CN03 area was determined. The results show a linear decrease in the concentration of SO$_4^{2-}$ with depth, ranging from 27.2 mM at 24 cmbsf to 2.7 mM at 290 cmbsf, concomitant with a gradual increase in the concentration of ΣH$_2$S from 1.2 to 6.28 mM for the same depth interval, respectively (Figure 2). However, the highest concentration of ΣH$_2$S was measured at 13.38 mM at 244 cmbsf. The methane flux was estimated to ~80 mmol/m$^2$/yr in 15GC, based on the linear zone in the sulfate concentration profile. The SMTZ in 15GC was estimated to be at ~205–255 cmbsf, which is supported by the peak at 244 cmbsf in the $^{31}$C$_{DIC}$ measurements (Chen et al., 2011). Consumption of sulfate is mainly ascribed to AOM (85%) and oxidation of other organic material (15%; Chen et al., 2011). A characteristic SMTZ was not observed in 29ROV. In 15GC, the methane concentration in the headspace of pore-water samples ranged from 0.012 to 0.38 mmol/L in horizons above the SMTZ, and gradually increased values from 0.16 to 3.6 mmol/L between 243 and 290 cmbsf (Vaulier, 2011).

The 15GC core consists of silt and clay rich sediments with a high bulk density (1.9–2.3 g/cm$^3$) and low porosity, presented as fractional porosity (~25–45%), and a low permeability (~30–60 cm/s). In comparison, bacterial 16S rRNA gene copies dominated 29ROV, with a high bulk density (1.9–2.3 g/cm$^3$) and low porosity, presented as fractional porosity (~25–45%), and a low permeability (~30–60 cm/s). The methane flux was estimated to ~80–90 cmbsf and ending at ~230–240 cmbsf, within the present SMTZ, is pierced by piping structures and chemosynthetic shells and shell fragments, indicating that the core site has been an active methane seeping area at earlier stage. The high variability in the carbonate content, represented by the XRF analysis on the Ca element, shows that that episodic activity of biogenic production is also found in more recent times, at a depth of ~30 cm (Figure 2).

**QUANTITATIVE PCR**

Both Bacteria and Archaea in 15GC were enumerated as 16S rRNA gene copies/g sediment using quantitative PCR. The relative abundance of 16S rRNA gene copies/g sediment throughout the core was 2.05 × 10$^{-6}$–1.06 × 10$^{-7}$ for Bacteria and 5.76 × 10$^{-6}$–5.75 × 10$^{-7}$ for Archaea (Figure 2). At all depths Archaea dominated over Bacteria, accounting for 51.7–93.3% of all 16S rRNA gene copies. The number of 16S rRNA gene copies of archaeal origin increased toward the SMTZ and decreased below this zone. In comparison, bacterial 16S rRNA gene copies dominated 29ROV in the horizons from the sediment surface to 7–8 cmbsf, ranging between 6.72 × 10$^{-6}$ and 9.1 × 10$^{-6}$ 16S rRNA gene copies/g sediment (Roalkvam et al., 2011). The bacterial population in 15GC was thus between two and three orders of magnitude lower than in 29ROV, except for the deepest horizon in 29ROV where the bacterial population decreased rapidly. Finally, the increase in the archaeal population with increasing depths was one order of magnitude for 15GC and three orders of magnitude for 29ROV.

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**Roalkvam et al.**

**ANME-stratification influenced by methane flux**

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In summary, the 16S rRNA gene quantifications showed two to three orders of magnitude lower relative cells numbers in the core from the micro-seepage CN03 area than in the active G11 pockmark.

TAXONOMY

The application of pyrosequencing of 16S rRNA gene tagged PCR-amplicons to obtain detailed knowledge on the community structure has recently been proven efficient in studying stratification of microorganisms in sediment cores at a much higher resolution than has been done previously (Lanzén et al., 2011; Roalkvam et al., 2011). This approach was used to examine the community structure in 15GC, from seven depth horizons (10, 30, 80, 120, 180, 240, and 270 cmbsf) analyzed by 454-pyrosequencing yielding 3344-26491 reads, whereof 16.4–29.0% were removed due to poor quality or chimeric sequences. The remaining number of reads in the dataset were between 2388 and 21400 for 15GC subsamples (Table 1).

Taxonomic classification revealed a high abundance of taxa that are deficiently described below phylum and class level, such as uncultivated taxa within Marine group 1 (MG-1; Thaumarchaeota); Thermoplasmata and MBG-B/DSAG (Crenarchaeota); Planctomycetes, Chloroflexi, and Candidate divisions (JS-1 and OP8; Bacteria). Hence, the number of classified reads at lower taxonomic levels decreased, where up to 88.6 and 89.8% of the community remained unclassified at order or family level, respectively. Therefore, only reads binned at phylum and class level, in addition to selected groups within Methanomicrobia (Euryarchaeota), were used further in this work.

Table 1 | Statistical parameters of 15GC.

| Depth (cmbsf) | Number of reads | Number of OTUs |
|-------------|----------------|----------------|
| 10          | 20471          | 455            |
| 30          | 18141          | 791            |
| 80          | 4859           | 144            |
| 120         | 2601           | 84             |
| 180         | 2388           | 107            |
| 240         | 17054          | 206            |
| 270\a       | 19501          | 131            |
| 270\b       | 20367          | 106            |
| 270\c       | 21400          | 138            |

\a\text{Amplicons from sample 270 cmbsf was made in triplicates.}

**FIGURE 2** The analyzes of geological parameters and physical properties in core 15GC show (from left to right): X-ray image of the core, Lithological log, Grain size, Bulk density, and fractional porosity based on MST logger unit, Ca, and S (count per second) based on XRF element core scan, Geochemical analyzes of H2S and SO4\textsuperscript{2−}, Enumeration of archaeal and bacterial 16S rRNA gene copies/g sediments based on quantitative PCR. Legends for Lithology: (...) silty-clay, (!) piping, and bioturbation, (ζ) shell/shell fragments, (♦) subsamples for DNA extraction, (+) subsamples for 16S rRNA gene tagged amplicon library construction.
MICROBIAL DIVERSITY IN AMPLICON LIBRARY AND ABUNDANCE

Most of the detected bacterial taxa were found in the shallower horizons, including phyla such as Proteobacteria, Planctomycetes, Deinococcus-Thermus, and the Candidate division OP8, all decreasing in abundance to less than 1% below horizons at 80–120 cmbsf (Figure 3A). Similarly, the abundance of the phylum Chloroflexi decreased rapidly in horizons below 80 cmbsf, although comprising up to 2% in some of these horizons. The Bacteria was dominated by the Candidate division JS-1, which was present throughout the sediments with the highest abundance at 10 and 30 cmbsf accounting for 13.2 and 28% of the total number of reads, respectively. In each sediment horizon, uncultivated lineages of Archaea dominated, congruent with the quantitative PCR-data (Figure 2). Low-abundance archaeal taxa, such as MCG (1.1–1.8%) and Group C3 (group 1.2; 1.2–2.6%) within Crenarchaeota and Thermoplasmata (0.3–4.2%) and Archaeoglobi (1.2%) within Euryarchaeota were mainly present in horizons between 10 and 80 cmbsf. Different depth profiles of the most abundant taxa MG-1, DSAG, and Methanomicrobia were observed (Figure 3A). The shallower horizons (10–120 cmbsf) above the SMTZ had high abundance of MG-1, ranging between 15.2 and 61.4%, which rapidly decreased toward the SMTZ (5.2% at 180 cmbsf and further to <1% in deeper parts of the core). The DSAG did not reach as high relative abundance as the MG-1, but represented a high share of the microbial community above and within the SMTZ, comprising between 16.2 and 29.2% in horizons at 10–240 cmbsf, except at 270 cmbsf where the abundance was reduced to 3.9% (Figure 3A). In the deeper horizons (180–270 cmbsf), where methane concentrations up to 3.6 mmol/L has been detected in the pore-water (Vaular, 2011), ANME clades affiliated with Methanomicrobia were increasingly dominant. A similar stratification of dominating ANME clades with increasing depth as in 29ROV was observed with a transition from an ANME-2a/b dominated community to an ANME-1 dominated community. However, the stratification was over wider sediment depths in 15GC ranging from 120 to 270 cmbsf in comparison to 4–22 cmbsf in 29ROV. The highest abundance of the ANME-2a/b clade was found at 120 and 180 cmbsf, with 16.9 and 33.3% of the total reads respectively (Figure 3B), and hence comprising a similar share of the community as in 29ROV (Roalkvam et al., 2011). The abundance of ANME-2a/b decreased gradually to <1% with increasing depth, while ANME-1 increased from 11.1

![Figure 3A](https://via.placeholder.com/150)

**FIGURE 3** | Microbial community structures at different depths in 15GC, based on 454-pyrosequencing of 16S rRNA gene tagged amplicons. The distribution of selected taxa is shown at phylum/class level (A) and within the class Methanomicrobia (B).
to 47.9% between 180 and 240 cmbsf. A further increase to 82.2% at 270 cmbsf was observed, which was the highest abundance of ANME-1 in 15GC (Figure 3B). The highest relative abundance of ANME-1 in 15GC (47.9–82.2%) corresponded to the abundance of ANME-1 in 29ROV (64–89%; Roalkvam et al., 2011). However, the abundance of ANME-2c increased to a maximum of 60% at 20–22 cmbsf in 29ROV (Roalkvam et al., 2011), which is considerably higher than the maximum value of 5.7% at 270 cmbsf in 15GC.

**DIVERSITY INDICES AND OTU DISTRIBUTION**

Shannon–Weaver index and Rao’s quadrate entropy index were used to evaluate and compare the microbial diversity of the communities in CN03 with those in the G11 pockmark, based on all reads in the amplicon libraries from 15GC and 29ROV. The main difference between the indices used is that Rao’s quadrate entropy index includes the distance between OTUs in addition to the abundance of sequences. A total of 3322 OTUs were obtained from 29ROV and 15GC combined, based on 193363 16S rRNA gene sequences from amplicon libraries. This approach revealed a vertical variation in microbial diversity in both cores, where a trend of decreasing diversity with increasing depth was observed regardless of the index used (Figure 4). For core 15GC, the diversity was decreasing gradually with depth, except at 120 cmbsf where the trend was interrupted by the particularly low diversity estimate. The core 29ROV had a different diversity profile, with a gradual decrease in the upper part of the core, followed by a considerable decline in the deeper part (Figure 4). The lowest diversity was found at 270 cmbsf in 15GC and 14–16 cmbsf in 29ROV, which corresponds to the horizon in each core with the highest abundance of ANME-1.

Sequences from 29ROV were clustered into 2370 OTUs (1908 unique OTUs) and 15GC were assigned to 1414 OTUs (952 unique OTUs sampling site) (Table 2), where only 462 OTUs were shared between the sampling sites. The majority of the taxa from both cores were clustered into several OTUs, where at least one OTU was common. Predominating taxa within each core were present in common OTUs, however some low-abundant taxa were present in one OTU (OTU_542; Figure 5), comprising 80.0–98.5% of the reads in 29ROV and 72.8–100% in 15GC (Table S1 in Supplementary Material). The remaining ANME-1 sequences were present in two additional OTUs, which also included sequences from the shallow horizons above the ANME-dominated zone in both cores. The ANME-2a/b affiliated sequences were mainly distributed in two dominating OTUs (Figure 5), one specific OTU for each core. OTU_442 comprised between 78.6 and 99.9% of ANME-2a/b sequences in 15GC, while OTU_50 constituted 54.5–95.2% of the sequences in 29ROV (0–10 cmbsf). Furthermore, some ANME-2a/b sequences from the ANME-1-dominated horizons in 29ROV were also assigned OTU_442.
Table 2 | Distribution of OTUs in 15GC and 29ROV.

| OTUs            | 15GC  | 29ROV |
|-----------------|-------|-------|
| Total number    | 1414  | 2370  |
| Number of unique| 952   | 1908  |
| Number of shared| 462   | 462   |
| Singletons      | 541   | 959   |
| Archaeal        | 160   | 267   |
| Bacterial       | 1242  | 2085  |
| Unassigned      | 12    | 18    |

Although the abundance of ANME-2c was considerably higher in 29ROV than 15GC, the majority of the reads were clustered into two common OTUs (OTU_168 and OTU_1280; Figure 5). In addition, a substantial number of reads from 29ROV were assigned to OTU_1800, which was unique for this core.

**DISCUSSION**

**METHANE FLUX AND STRATIFICATION OF METHANOTROPHS**

The Nyegga area is characterized by numerous pockmarks and methane seepage structures in different developmental stages indicating a dynamic, temporal, and spatial system where different geochemical settings may influence the microbial community structure. In this study, 454-pyrosequencing of 16S rRNA genes tagged amplicons were used to compare the microbial stratification in a gravity core (15GC) from the less active CN03 area with that in a push core (29ROV) from the active seepage structure G11 pockmark. The CN03 area is characterized by micro-seepage of methane, few pockmarks, a SMTZ located at ~200–250 cmbsf and a BSR at 250–300 mbsf. The G11 pockmark in comparison, is characterized by shallow gas hydrates and SMTZ, authigenic carbonates, and pingo structures within the pockmark. The higher methane flux sustains macro-fauna and bacterial mats (Hovland and Svensen, 2006; Chen et al., 2010; Ivanov et al., 2010). The methane fluid flux was ~80 mmol/m² year for 15GC and 300–540 mmol/m² year for 29ROV (Chen et al., 2010). The methane flux in the CN03 area is within the same range as other seep areas where ANME have been found to be abundant, such as the Santa Barbara Basin (164–200 mmol/m² year; Harrison et al., 2009) and Gulf of Mexico (20–200 mmol/m² year; Coffin et al., 2008; Lloyd et al., 2010). In 29ROV, the flux is apparently so high that methane reaches the sediment surface where it stimulates a dominance of aerobic methanotrophic Gammaproteobacteria (Roalkvam et al., 2011). In 15GC, aerobic methanotrophic Gammaproteobacteria were not identified, indicating that the methane seeping through the sediments were consumed by ANME which were increasingly dominating the amplicon libraries from 120 cmbsf and below (Figure 3). A dominance of ANME in deeper horizons was also observed in 29ROV, however, the ANME-dominated sediment horizons extended over a wider depth interval in 15GC. Furthermore, similar relative abundances and equivalent transitions between ANME-2a/b and ANME-1 with increasing depth (Figure 3B) were observed in both 29ROV and 15GC cores. However, the proportion of ANME-2c seemed to differ between the cores as 60% of all reads were assigned to ANME-2c in 29ROV, whereas only 5.7% of all reads were assigned to this taxonomic group in 15GC. It is possible that the observed difference in ANME-2c abundances is an effect of the much wider methane gradient in 15GC and that ANME-2c at this sampling site are found in higher numbers in horizons deeper than 270 cmbsf, and thus not detected in our study.

The methane flux seems to have little effect on the stratification of the ANME groups 2a/b, 1, and 2c in the Nyegga field, as a similar stratification of these clades was found in both the 29ROV (Roalkvam et al., 2011) and 15GC locations (Figure 3B). This suggests that shifts in ANME clades through the cores are determined by other factors than methane availability. Moreover, both cores are dominated by the same OTUs of ANME-1 and ANME-2c, indicating that organisms potentially adapted to different methane fluxes cannot be distinguished on the OTU level within these clades. On the other hand, differences in methane flux may partly explain why different OTUs of ANME-2a/b dominate in the cores. The methane flux seems to largely influence the specific density of ANME in the cores as the number of archaeal 16S rRNA genes per gram of sediment in the ANME-dominated horizons was observed to be two orders of magnitude lower in 15GC (Figure 2) than in 29ROV (Roalkvam et al., 2011). This is congruent with previous studies of sediments with variations in methane flux and depth of the SMTZ where ANME comprise up to $3 \times 10^9$ cells/cm² sediment in marine sediments with shallow gas hydrates, such as Gulf of Mexico (Orcutt et al., 2008), Hydrate Ridge (Knittel et al., 2005), and Eckernförde Bay (Treude et al., 2005a), whereas in low methane seepage areas with deeper gas hydrates, such as Santa Barbara and Peru Margin, ANME are rare or absent (Biddle et al., 2006; Inagaki et al., 2006; Harrison et al., 2009).

Both in the sediments from the G11 pockmark and in 15GC, ANME-2a/b was found to dominate in horizons with lower concentrations of sulfide compared to the ANME-1 dominated horizons. These results are in accordance with previous studies suggesting that ANME-2 is sensitive to H₂S produced during AOM with sulfate (Meulepas et al., 2009a,b). Furthermore, the abundance of ANME-2a/b was found to be negatively correlated with high methane and sulfide concentrations in Guaymas Basin sediments, whereas an opposite correlation was found for ANME-1 (Biddle et al., 2011). Also, a zonation of ANME-2 communities in areas with efficient H₂S removal and ANME-1 in zones with higher concentrations of H₂S was observed in multilayered microbial mats in the Black Sea (Krüger et al., 2008). Due to the higher methane flux in G11 the ANME-2a/b population is exposed to methane-rich fluids whereas in CN03, the dominating ANME-2a/b is found in horizons with low methane concentrations of 0.035–0.13 mmol/L. However, the piping structures and shell fragments in these horizons show that the methane concentration has probably been higher in earlier times, indicating that the ANME-2a/b population could have been established in a methane-enriched environment in the past and that the ANME-2a/b population present in 15GC now is sustained by lower methane concentrations. Given the ability to survive in methane depleted environments, ANME-2a/b could be the seed population in new methane-enriched systems, as suggested by Knittel and Boetius (2009). Thus, in order to fully understand the stratification and dynamics of ANME...
communities at Nyegga, more knowledge on temporal variations in the microbial community structures might be needed.

SYNTROPHIC PARTNERS OF ANME

The AOM with sulfate has in previous studies been shown to be performed by ANME in syntrophy with sulfate-reducing Deltaproteobacteria, where ANME-1 and ANME-2 are mainly associated with sulfate-reducers within Desulfosarcina and Desulfococcus (DSS; Knittel et al., 2005; Schreiber et al., 2010), while ANME-3 are associated with sulfate-reducers within Desulfobulbus (DBB; Niemann et al., 2006; Lösekann et al., 2007). The share of Deltaproteobacteria in horizons with the highest relative abundance of ANME-2a/b was ranging between 9.0 and 9.1% in 29ROV (Roalkvam et al., 2011), but was below 0.7% in
15GC (Figure 3A) of which most sequences were assigned to a clade that is not assumed to be a syntrophic partner of ANME (Table S1 in Supplementary Material). From the low-abundance of detected Deltaproteobacteria, it was not obvious to us which of the detected organisms that acted as a sulfate-reducing syntrophic partner for ANME-2a/b, at least not in 15GC. One possibility is that the abundance of Deltaproteobacteria is underestimated due to bias in the PCR amplification of 16S rRNA genes. Another possibility is that other organisms than Deltaproteobacteria act as the sulfate-reducing syntrophic partner for the ANME-2a/b detected in the Nyegga field. JS-1 was present in all ANME-2 dominated horizons in 15GC, but was even more dominating in other sediment horizons, indicating no obligate relationship with ANME. The JS-1 group is ubiquitous in marine sediments, at depths ranging from <10 to >200 mbsf (Rochelle et al., 1994; Inagaki et al., 2006; Parkes et al., 2007; Webster et al., 2007). Inagaki et al. (2006) hypothesized that JS-1 could be adapted to anaerobic conditions in organic-rich sediments associated with methane hydrates, which is similar to the environment at Nyegga. Further indications of JS-1 being a heterotrophic sulfate-reducing bacterium are based on enrichment cultures where sulfate was depleted in wells with acetate (Webster et al., 2011). In 29ROV, the abundance of JS-1 throughout the core was correlated to the abundance of ANME-2 (Roalkvam et al., 2011), and ANME-2 may benefit from the activity of JS-1 by transfer of reducing equivalents derived through AOM (Roalkvam et al., 2011), which corresponds well with the average TOC values (~0.5–1%) for the region (Hölemann and Henrich, 1994). In sedimentary environments, the organic matter buried over geological timescales is utilized as an energy source by organotrophs (Kujawinski, 2011; Orcutt et al., 2011). Through microbial remineralization, degradation products are formed which can be utilized by diverse heterotrophic taxa. The microbial distribution in 15GC showed a high abundance of taxa that generally occurs in high numbers in marine sediments, such as Planctomycetes, Chloroflexi, Bacteroidetes, JS-1, MBG-B/DSAG, MG-1, MCG, and MBG-D (Figure 3A; Reed et al., 2002; Inagaki et al., 2006; Harrison et al., 2009; Blazjak and Schippers, 2010). In 15GC, Archaea outnumbered Bacteria in all horizons (Figure 2) with a clear stratification of the archaeal phyla (Figure 3A). The MG-1 predominated in horizons probably depleted in oxygen at 10–120 cmbsf. Cultivated representatives of MG-1 have been shown to perform aerobic ammonium oxidation (Hallam et al., 2006; Nicol and Schleper, 2006; Walker et al., 2010). Ammonium, potentially derived from the degradation of nitrogen-containing organic matter, could support the population of MG-1 in 15GC and may possibly be oxidized anaerobically. The MG-1 is a diverse clade with several subgroups that are not well characterized (Durbin and Teske, 2010), and the MG-1 could possibly have a wider range of useful metabolisms in this environment. Within the Archaea, the DSAG is predominant in several marine environments, such as deep sea sediments (Vetriani et al., 1999; Fry et al., 2008; Wang et al., 2010), sediments overlaying shallow gas hydrates (Inagaki et al., 2006) and within the SMTZ at Santa Barbara Basin (Harison et al., 2009) and Peru Margin (Sorensen and Teske, 2006). In 15GC, DSAG was uniformly distributed in horizons between 10 and 240 cmbsf where the concentration of methane is low, but was outcompeted by ANME-1 at 270 cmbsf where the concentration of methane is higher. Hence, DSAG may rather perform organotrophic sulfate reduction as suggested by Biddle et al. (2006) and Inagaki et al. (2006) than consumption of methane in 15GC. This is also supported by the low-abundance of DSAG in 29ROV, where the horizons are exposed to methane-rich fluids. The higher methane flux in the G111 porewater apparently also influenced the absolute numbers of microorganisms, with two to three orders of magnitude higher number of 16S rRNA gene copies per gram of sediment compared to the CN03 area. Hence, even though taxa such as Planctomycetes, Chloroflexi JS-1, MG-1, and DSAG were present with abundances between <1 and 10% in 29ROV, their absolute numbers are in the same order of magnitude as in the 15GC core as the total cell number in these horizons were two to three order of magnitudes higher than in the 15GC. This indicates that both sampling sites have equivalent amounts of microorganisms potentially involved in degradation of organic matter, which is consistent with the relatively even distribution of organic matter in Nyegga sediments (Hölemann and Henrich, 1994; Ivanov et al., 2010).
CONCLUSION
In conclusion, our results show that ANME dominate in Nyegga cold seeps, both in the micro-seeping CN03 area and the highly active G11 pockmark. An equivalent stratification of ANME clades was observed at both sampling sites, indicating that the vertical distribution of ANME is influenced by increasing sulfide concentration in the sediments or other factors rather than the methane fluid flux. Furthermore, a fine-scale view into the ANME communities at each location was achieved through OTU clustering of ANME-affiliated sequences from the amplicon libraries, revealing a common OTU of ANME-1, and unique OTUs of ANME-2a/b at the sampling sites. Based on our results we suggest that free-living ANME-1 is the most abundant taxon, and also the main consumer of methane in Nyegga cold seeps which can reduce the methane emission to the atmosphere from sources within the sediments. The specific ANME-2a/b OTUs could reflect adaptations to the geochemical composition at each location, with different affinities to methane. The ability to survive in marine systems depleted in methane could also be advantageous for potential seed populations.

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SUPPLEMENTARY MATERIAL
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