Detection of putatively thermophilic anaerobic methanotrophs (ANMEs) in diffuse hydrothermal vent fluids

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Abstract

The anaerobic oxidation of methane (AOM) is carried out by a globally distributed group of uncultivated Euryarchaeota, the anaerobic methanotrophic archaea (ANME). In this work, we used G+C analysis of 16S rRNA genes to identify a putatively thermophilic ANME group and applied newly designed primers to study its distribution in low-temperature diffuse vent fluids from deep-sea hydrothermal vents. We found that the G+C content of the 16S rRNA genes ($P_{GC}$) is significantly higher in the ANME-1GBa group than in other ANME groups. Based on the positive correlation between the $P_{GC}$ and optimal growth temperatures ($T_{opt}$) of archaea, we hypothesize that the ANME-1GBa group is adapted to thrive at high temperatures. We designed specific 16S rRNA gene-targeted primers for the ANME-1 cluster to detect all phylogenetic groups within this cluster including the deeply branching ANME-1GBa group. The primers were successfully tested both in silico and in experiments with sediment samples where ANME-1 phylotypes had previously been detected. The primers were further used to screen for the ANME-1 microorganisms in diffuse vent fluid samples from deep-sea hydrothermal vents in the Pacific Ocean, and sequences belonging to the ANME-1 cluster were detected in four individual vents. Phylotypes belonging to the ANME-1GBa group dominated in clone libraries from three of these vents. Our findings provide evidence of existence of a putatively extremely thermophilic group of methanotrophic archaea that occur in geographically and geologically distinct marine hydrothermal habitats.

Introduction

Over the past decade, many studies on the anaerobic oxidation of methane (AOM) have been published (for a review see (1)), emphasizing the global distribution of this process in marine
sediments and its importance as a factor that reduces the level of methane emissions to the atmosphere. The oxidation of methane by archaea is assumed to be a reverse methanogenesis coupled to the reduction of sulfate by sulfate-reducing bacteria (2). This process occurs in marine habitats where sulfate from the seawater and methane of biological and geochemical origin from deeper layers meet (1). Based on the 16S rRNA phylogeny, all anaerobic methanotrophic archaea (ANME) are grouped into three distinct clusters of Euryarchaeota, namely ANME-1, ANME-2, and ANME-3 (3, 4). Microorganisms of the ANME-2 and ANME-3 clusters belong to Methanosarcinales whereas ANME-1 cluster is a deep phylogenetic branch of Euryarchaeota only distantly related to the orders Methanomicrobiales and Methanosarcinales (5). Despite the great interest in these microorganisms, all three clusters remain uncultured. It is assumed that one of the major obstacles to the isolation of AOM-mediating microorganisms is their slow growth, the main reason for which is presumably bioenergetic limitations caused by the very low energy yield of AOM (6). According to theoretical calculations, the free-energy yield (∆G) of sulfate-dependent oxidation of methane increases with increasing temperature (7), suggesting that high-temperature AOM should occur. There are several lines of evidence that anaerobic methanotrophs can thrive at high temperatures. Sequences of the 16S rRNA genes of all three clusters of methane oxidizing archaea (ANME-1, ANME-2 and ANME-3) as well as ANME-specific core lipids have been found in deep-sea hydrothermal vents and warm sediments (8-14). Moreover, several researchers managed to determine in-vitro methane oxidation rates at different temperatures by using hydrothermal sediment samples from Guaymas Basin and Middle Valley (7, 15, 16). These studies have shown maximum AOM activity between 45 °C to 60 °C. Here, we provide one more line of evidence of thermophilic AOM by analysis of the G+C content of the 16S rRNA genes (hereafter P_{GC}) of ANME phylotypes and application of newly designed specific primers.

The positive correlation between the G+C content of the ribosomal RNA and optimal growth
temperatures of prokaryotes (hereafter $T_{\text{opt}}$) was noted by the founders of 16S rRNA-based phylogeny (see, e.g., (17)). The relevant data were systematized by Galtier and Lobry (18) and further formalized by Kimura et al. (19), who obtained the correlation equation $P_{GC} = 0.17T_{\text{opt}} + 49.0$ ($R^2=0.828$) but mentioned occurrence of deviations from the general trend. These deviations are especially notable in the domain Bacteria. In the present work, we show that, in Archaea, the correlation is stringent enough to allow confident predictions of the thermophilic adaptation of uncultured microorganisms with $P_{GC}$ higher than a particular value. We identify a particular phylogenetic group of methane oxidizing archaea with a high $P_{GC}$ as putatively thermophilic. We then designed, tested, and applied a specific primer set to assess distribution of putatively thermophilic anaerobic methanotrophs in geographically and geologically distinct deep-sea hydrothermal vent fluids.

Materials and Methods

Analysis of the correlation between the $P_{GC}$ of archaea and their $T_{\text{opt}}$. The list of the currently recognized archaeal species (a total of 374) and their type strains was taken from the J.P. Euzéby's LPSN site (http://www.bacterio.cict.fr/; 20) (for final check, update of April 13, 2012 was used). The 16S rRNA gene sequences were downloaded from RDP Release 10 (http://rdp.cme.msu.edu/) or from GenBank following links at the Euzéby's site. The G+C content was determined using BioEdit 7.0.9.0 program (21). The $T_{\text{opt}}$ were taken from relevant original publications. For 19 type strains, the sequences were not available or were too short ($<1200$ nt) or contained introns or $T_{\text{opt}}$ could not be found. Thus, the correlation was analyzed for a total of 355 archaeal type strains.

The $P_{GC}$ was also determined for all archaeal isolates with 16S rRNA gene sequences longer...
than 1200 nt available from RDP Release 10 (for final check, Update 28, Jan 12, 2012 was used, 2022 sequences). Then, we examined the information available on those of the isolates that were not affiliated with known thermophilic genera but exhibited \( P_{GC} \) higher than a particular value (see Results and discussion).

**Analysis of the \( P_{GC} \) of ANME phylotypes.** 945 sequences of 16S rRNA genes affiliated with five ANME groups (ANME-1a, ANME-1b, ANME-2ab, ANME-2c and ANME-3) were downloaded from the ARB-SILVA database (http://www.arb-silva.de) (22). Eight sequences affiliated with the ANME-1AT and ANME-1GBa groups were downloaded from the database developed by German Jurgens (5). Five more sequences affiliated with the ANME-1GBa group were taken from GenBank by accession numbers in (14). The G+C content of ANME phylotypes was determined using BioEdit 7.0.9.0 program.

**Design and specificity verification of primers targeting the ANME-1 cluster.** For the design of the ANME-1-specific primers, we retrieved 275 sequences affiliated with the ANME-1a and ANME-1b groups from the ARB-SILVA database and 8 sequences affiliated with the ANME-1AT and ANME-1GBa groups from the database developed by German Jurgens (5). The primers were designed using home-made software briefly described in (23) with weighting mismatches as described in (24). At the stage of primer design, a restricted outgroup sampling was used. However, the specificity of the primers was further confirmed by using the RDP Probe Match tool and the OligoReport program (K.E. Ashelford, unpublished) with the RDP database (http://rdp.cme.msu.edu/) for individual primers and by using the OligoCheck program (K.E. Ashelford, unpublished) with the ARB-SILVA database for primer pairs. The newly designed primers, as well as the previously published primers and probes targeting phylotypes of the ANME-
Field Sites and Sampling. Hydrothermally heated sediments characterize the Guaymas Basin hydrothermal vent site in the Gulf of California. In 2010, during the research cruise BIG, a core sample of sediments from Guaymas Basin was obtained (location BIG 1). Sediments in the sampling area were covered with a white microbial mat. For DNA extraction, the layer of 4 to 10 cm below the sediment surface was used. The temperature in this layer ranged from 50 °C to 70 °C.

Mississippi Canyon Block 118 (MC118) in the Gulf of Mexico is characterized by methane hydrate deposits and thermogenic hydrocarbon-rich fluids (30). It is located offshore of Louisiana at a water depth of ~890 m. Samples of sediments covered with a white microbial mat were taken in 2006 using the Johnson-Sea-Link submersible. Temperature of bottom water was 5.5 °C. Detailed description of sediments of the Mississippi Canyon 118 (MC118) in the Gulf of Mexico is provided in (30, 31).

From 2006 through 2009, vent fluids were collected from seven hydrothermal sites in the Pacific Ocean: Axial Seamount and the Endeavour Segment (32), both on the Juan de Fuca Ridge, and five volcanoes along the Mariana Arc (33) (Supplemental Material Fig. A1, Table A1). All fluid samples were collected from low-temperature vents using the Hydrothermal Fluid and Particle Sampler (HFPS) (34) mounted on the deep-sea research submersibles Jason 2 and Alvin. While on the seafloor, fluids were passed through a Sterivex-GP (0.22-μm pore size). These filters were immediately frozen at -80°C upon vehicle recovery.

Characteristics of all samples that produced positive PCR results with the newly designed primer sets are shown in Table 2.

Nucleic acid extraction. DNA from vent fluid filters was extracted as previously described
(35). DNA from a sediment core of a Guaymas Basin hydrothermal field was extracted as previously described (36). DNA of the sediments of the MC 118 site was kindly provided by the Andreas Teske Laboratory (University of North Carolina at Chapel Hill).

**PCR amplification of 16S rRNA gene fragments.** PCR amplification was performed in a 50-μl reaction mixture containing 1 μl of template DNA, 1× GoTaq Green Reaction Buffer (Promega, San Luis Obispo, California), 1 μl of 10 mM 4dNTP mix (200 μM final concentration for each dNTP) (Promega, San Luis Obispo, California) and 1 U of GoTaq DNA Polymerase (Promega, San Luis Obispo, California). Concentrations of the primers were calculated using the «PCR Optimization» program (http://molbiol.ru/eng/scripts/01_14.html). PCR programs were adjusted experimentally based on the standard PCR protocol (37) and on the calculations of the annealing temperatures of primers made using the OligoAnalyzer program (http://eu.idtdna.com/analyzer/Applications/OligoAnalyzer/). For the primer set ANME-1-25(F) – ARCH-915(R), the annealing temperature was 59 °C, and the concentration of each primer was 0.45 μM. For the primer set ANME-1-25(F) – ANME-1-1406(R) the annealing temperature was 58°C and the concentration of each primer was 0.3 μM. For the primer set ANME-1-1118(F) – ANME-1-1406(R) the annealing temperature was 61°C and the concentration of each primer was 1.25 μM. As negative controls, reactions without added DNA, reactions with the addition of DNA isolated from the surrounding sea water, and reactions with the addition of genomic DNA of *Methanocaldococcus jannaschii* or *Archaeoglobus profundus* were used. As positive controls, reactions with the addition of plasmid DNA harboring a cloned ANME-1 16S rRNA gene fragment (kindly provided by the Teske Laboratory (University of North Carolina at Chapel Hill)) were used. Polymerase chain reactions were performed using a Mastercycler Gradient (Eppendorf, Hamburg, Germany). Amplicons were visualized with ethidium bromide on 1% agarose gels in 1×TAE buffer.
Cloning and sequencing of PCR-amplified 16S rRNA gene fragments. PCR products were purified and concentrated using the MinElute PCR Purification Kit (Qiagen) according to the manufacturer’s instructions. Product quality was assessed on 0.8% agarose gels stained with ethidium bromide. Bands were excised and DNA was extracted using the MinElute Gel Extraction Kit (Qiagen). This purified product was ligated into pCR4-TOPO vector for 5 min at room temperature and transformed into electrocompetent cells according to the manufacturer’s instructions (Invitrogen). For each library, 24-96 clones were randomly selected and grown in SuperBroth with 50 mg ml⁻¹ kanamycin in 96 deep-well blocks overnight at 37°C with vigorous shaking. Cells were collected by centrifugation and plasmid DNA was isolated using a standard alkaline-lysis procedure (38). Plasmids were sequenced bidirectionally with primers T3 (5′-ATTACCCTCACTAAAGGGA) and T7 (5′-TAATACGACTACTATAGGG) using Big Dye Terminator v.3.1 kit on an ABI 3730 sequencer (Applied Biosystems, USA). If necessary, the intermediate primer ARCH-915(R) (29) was used.

Sequence analysis. Sequences were analyzed and edited in the Chromas Lite 2.01 program (http://www.technelysium.com.au). Forward and reverse reads were assembled into contigs using the BioEdit 7.0.9.0 program. Sequences were aligned in the ClustalW program (39). All sequences were analyzed by the Pintail program (40) in order to detect chimeric 16S rRNA gene sequences. Sequences originating from the same samples were grouped into 98% similar operational taxonomic units (OTUs) using the cd-hit program (41). The phylogenetic tree (Fig. 3) was constructed with the ARB software package (42) using maximum-likelihood (PHYML) algorithm and non-parametric bootstrap analysis (100 replicates). The sequences obtained in this study were deposited in NCBI GenBank with accession numbers JQ740748-JQ740762.
results and discussion

correlation between the $P_{GC}$ of archaea and their $T_{opt}$. We plotted the G+C contents of 355 16S rRNA genes of archaean type strains against their optimal growth temperatures. Our data can be approximated by the linear correlation equation $P_{GC} = 0.18T_{opt} + 49.8; R^2 = 0.819$ (Fig. 1), which is close to the Kimura’s et al. equation (19), obtained for a less strictly defined organism sampling. Since G:C pairs are more stable than A:U pairs due to an additional hydrogen bond (43), we interpret the correlation as indicative of the necessity for the 16S rRNA molecule to have a definite rigidity level of its secondary structure.

The distribution of the dots in the plot in Fig. 1 shows that there exist particular $P_{GC}$ ranges characteristic of mesophilic ($T_{opt} < 50^\circ\text{C}$), moderately thermophilic ($50^\circ\text{C} \leq T_{opt} < 70^\circ\text{C}$), extremely thermophilic ($70^\circ\text{C} \leq T_{opt} < 80^\circ\text{C}$), and hyperthermophilic ($T_{opt} \geq 80^\circ\text{C}$) groups of archaean type strains. For the present work, the most significant conclusion from Fig. 1 is that all archaean type strains with $P_{GC}$ above 60 mol% are thermophiles and those with $P_{GC}$ above 63 mol% are extreme thermophiles or hyperthermophiles. Moreover, our analysis of all sequences of the 16S rRNA genes of archaean isolates available from the RDP database with checking the $T_{opt}$ of the corresponding microorganisms in relevant publications showed that no archaean isolates with a $P_{GC}$ above 60 mol% have been reported to be mesophilic. The information on the four isolates that had a $P_{GC}$ above 60 mol% and were not affiliated with known thermophilic genera is presented in Supplemental Material Table A2 (these are haloarchaean with unreported temperature optima). It follows that a $P_{GC}$ value exceeding 60 mol% may be accepted as sound evidence of the thermophilic nature of the analyzed archaean phylotypes.
Analysis of $P_{GC}$ of ANME phylotypes. We have analyzed the G+C content of over 900 sequences of 16S rRNA genes of seven ANME groups: ANME-1a (8), ANME-1b (4), ANME-1AT (5), ANME-1GBa (5), ANME-2ab (3), ANME-2c (3) and ANME-3 (4) (Fig. 2). We found that for six ANME groups (ANME-1a, ANME-1b, ANME-1AT, ANME-2ab, ANME-2c and ANME-3), $P_{GC}$ does not exceed 59 mol%. Meanwhile, the minimum value of this parameter for the group ANME-1GBa is 62.5 mol% and the average value is 63.5±0.7 mol%.

We also took a separate examination of the G+C contents of 16S rRNA sequences representing ANME microorganisms that were detected in or enriched from different hydrothermal-associated environments as some of such sequences were not in Silva database or do not have a defined phylogenetic affiliation there. For this analysis, 76 16S rRNA gene sequences representing all known ANME clusters and subgroups were taken from relevant publications (Supplemental Material Table A3). The $P_{GC}$ of these sequences was compared to corresponding in situ or incubation temperatures. All ANME sequences except those that belong to the ANME-1GBa group have $P_{GC}$ lower than 59 mol%. Based on the obtained correlation between the $P_{GC}$ of archaea and their $T_{opt}$, we can predict that these sequences may represent moderate thermophilic or mesophilic microorganisms, which is consistent with in situ or incubation temperatures reported in corresponding publications.

Based on our 60 mol% cut-off value, only microorganisms of the ANME-1GBa group can confidently be predicted to be adapted to thrive at a high temperature. In addition, considering $P_{GC}$ of other ANME groups and absence of extreme thermophiles or hyperthermophiles with $P_{GC}$ below 60 mol% in our plot (Fig. 1), we may state that, among the currently recognized methanotrophic archaea (Fig. 2), microorganisms of the ANME-1GBa group are the only candidates for extreme thermophily ($T_{opt} \geq 70 ^{\circ}C$). Previously, phylotypes of ANME-1GBa group were found only in hydrothermal sediments of the Guaymas Basin (8, 14). The ANME-1GBa group is the deepest
branch within the ANME-1 cluster and is phylogenetically distant from other members of this cluster (5). We failed to find published ANME-1-targeted primers that would be able to amplify all groups within this cluster, including the ANME-1G Ba group (Table 1). Therefore, we developed a primer system with the goal of screening for ANME-1G Ba phylotypes and other phylotypes of the ANME-1 cluster in low-temperature fluids from deep-sea hydrothermal vents. We rejected the idea of developing ANME-1G Ba specific primers because such primers would have to be based on a small number of highly similar sequences, and thus there was a risk of over-specificity to particular sequences.

**In silico verification of the specificity of the ANME-1-targeted primers.** Three ANME-1-targeted primers that we designed perfectly matched most of the sequences of all four phylogenetic groups within the ANME-1 cluster (Table 1). These primers are ANME-1-25(F) (5’-GAGCCYACTGCYATCAGMGT-3’), ANME-1-1118(F) (5’-CYCRCAGTTGCCAGCATCTC-3’), and ANME-1-1406(R) (5’-AYCYCACTCGGYTGCTTAGGAT-3’). Table 1 presents the results of our analysis of the specificity of the newly designed primers, as well as of previously published primers and probes. In contrast to our primers, none of the previously published primers and probes covered the ANME-1 cluster entirely, as expected since these primers were designed to target specific subsets of ANME-1 clusters.

All previously published and newly designed primers and probes were also checked *in silico* by using 16S rRNA gene sequences of archaeal type strains as negative controls (Table 1). Moreover, the specificity of the newly designed systems of primers was confirmed by using all archaeal 16S rRNA gene sequences (of cultured and uncultured archaea) from the ARB-SILVA database (more than 25,000 sequences). This analysis showed that the primer pairs ANME-1_25F - ANME-1_1406R and ANME-1_1118F - ANME-1_1406R did not match any sequences except for
sequences of ANME-1 group, even if one mismatch per primer was allowed. All three new primers did not match any bacterial 16S rRNA gene sequences. 

*In silico* analysis showed that the designed systems of primers do not form homo- or heterodimers with significant $\Delta G^0$. The pair of primers ANME-1-25(F) - ANME-1-1406(R) allows amplification of nearly entire 16S rRNA gene, while ANME-1-1118(F) - ANME-1-1406(R) primer pair will be suitable for applications that require a smaller size amplicon (~300 bp).

**Verification of the specificity of the newly designed primers in experiments with environmental samples.** The specificity of the designed primers was also confirmed in experiments with sediment samples where ANME-1 phylotypes had previously been detected: those from the hydrocarbon seeps in the Gulf of Mexico (MC118 site, (31)) and the hydrothermal sediments of the Guaymas Basin (8, 14).

Three sets of primers were tested with the sample from the Gulf of Mexico: ANME-1-25(F) - ANME-1-1406(R), ANME-1-1118(F) - ANME-1-1406(R) and ANME-1-25(F) – ARCH-915(R) (Table 2, Fig. 3). Using two different combinations of primers (ANME-1-25(F) - ANME-1-1406(R) and ANME-1-25(F) – ARCH-915(R)), sequences related to ANME-1a and ANME-1b were detected. In terms of the diversity of sequences and their ratio in clone libraries, the results of using these two systems of primers were nearly identical. However, only ANME-1b group was detected by using third primer set (ANME-1-1118(F) - ANME-1-1406(R)) (not shown in Fig. 2), in accordance with the reduced *in silico* specificity of the ANME-1-1118(F) primer to the ANME-1a group (Table 1). Previous work detected many different uncultured archaeal lineages in the MC118 site (31), but only the 16S rRNA genes of the ANME-1 cluster were detected with our new primers. The primer set ANME-1-25(F) – ARCH-915(R), in which the reverse primer is universal for archaea, also showed high specificity to the ANME-1 cluster.
The ANME-1-25(F) - ANME-1-1406(R) primer set was also verified by testing it with hydrothermal sediments from Guaymas Basin (site BIG 1). A broad diversity of ANME groups was previously detected in Guaymas Basin sediments, including two distinct offshoots of the ANME-1 cluster: ANME-1AT and ANME-1GBa (8). Both of these groups were detected using ANME-1-25(F) - ANME-1-1406(R) primer with sediment sample of site BIG 1 (Table 1; Fig. 3).

Detection of phylotypes of the ANME-1 cluster in diffuse vent fluids. In contrast to focused fluids that circulate through the oceanic crust at high temperatures (generally 250° to 350°C), diffuse flows provide large stable habitats beneath the seafloor with gradient of both temperature and redox pairs suitable for the development of microbial communities (44-46). In order to study the distribution of microorganisms of ANME-1GBa group as well as of other groups of ANME-1 cluster in diffuse hydrothermal ecosystems, we screened 50 hydrothermal vent fluid samples representing 42 different diffuse flow hydrothermal vents from the Axial Seamount, the Endeavour Segment, and from the volcanoes of the Mariana Arc for the 16S genes of the ANME-1 cluster using ANME-1-25(F) – ARCH-915(R) primer set. A positive PCR result was obtained for four samples. Amplified gene fragments were cloned and sequenced (Table 2).

Axial Volcano is an active submarine volcano located on the Juan de Fuca Ridge (47). Here we detected anaerobic methanotrophs of the ANME-1 cluster at Bag City vent, a stable, basalt-hosted diffuse vent located in lobate lavas (35). All ANME-1 microorganisms detected in this site were classified as members of ANME-1GBa group (Table 2; Fig. 3).

16S rRNA genes of microorganisms of ANME-1 cluster were also amplified from two vent fluid samples of Endeavour Segment – Easter Island and Boardwalk (32). In both cases, samples were taken in an area of primarily flat basalt with diffuse venting out of cracks, near high-temperature black smokers. All sequences obtained from Easter Island and the majority of sequences
obtained from Boardwalk were also related to the ANME-1GBa group (Table 2; Fig. 3).

Finally, microorganisms of ANME-1 group were detected at Fault Shrimp, a diffuse vent located at NW Rota-1, an actively erupting volcano on the Mariana Arc (48). Fault Shrimp has venting out of cracks in basaltic andesite with microbial mat and shrimp present. All ANME-1 microorganisms detected in this site were classified as members of ANME-1a group (Table 2; Fig. 3).

Like the sequences reported in previous works (8, 14), the ANME-1GBa OTUs that we detected at deep-sea hydrothermal vents had a significantly higher G+C content (62.1-64.5 mol%) than the OTUs of other ANME-1 groups. Taking into account the sequences obtained in our study, the PGC of ANME-1GBa group is 63.6±0.8 mol%. Considering the obtained correlation (Fig. 1) and the fact that there are no mesophilic or moderately thermophilic archaeal type strains with a PGC above 63 mol%, we predict that the optimal growth temperatures of microorganisms of ANME-1GBa group is above 70°C. The putatively thermophilic methanotrophic archaea of the ANME-1GBa group were previously detected only in hydrothermal sediments in the Guaymas Basin (8, 14).

Our findings provide evidence for wider distribution of this group in geographically and geologically distinct hydrothermal habitats. Moreover, ANME-1GBa group appears to be the most frequent subgroup of ANME-1 cluster in diffuse hydrothermal vent fluids. It should be noted, that phylotypes of ANME-1GBa group have only been found in environments associated with hydrothermal activity (hydrothermal sediments of Guaymas Basin and diffuse hydrothermal ecosystems). The same is true for mcrA genes that may be assigned to ANME-1GBa group (see below).

Microorganisms of ANME-1 cluster were detected in only 4 out of the 42 unsedimented basalt-hosted deep-sea hydrothermal vents studied. This suggests only sporadic occurrence of these microorganisms in such environments. Previous work indicates that non-sedimented environments
are not typical habitats for methanotrophic archaea. This can be explained by the absence of large sources of methane in such habitats. However, while there are no visible sediments at the surface of the studied sites of hydrothermal venting, chemical data suggests that there are large deposits of buried organic-rich sediments beneath the ridge crest at the Endeavour Segment (49-51). Here, due to the putatively buried sediments, methane levels in hydrothermal fluids are extremely high (up to 3 mM) suggesting a large source of methane for methanotrophy to occur in the subseafloor. There is no evidence for such sediments at Axial or NW Rota-1, but at each site, subsets of diffuse vents do have micromolar concentrations of methane, likely due to methanogenesis (32, 33). Taking into account (i) evidence of thermophilic nature of the ANME-1GBa group, (ii) low temperature of the studied fluids at the surface, (iii) absence of sediments at the surface of the studied sites and (iv) evidence of the presence of organic-rich sediments beneath the surface of Easter Island and Boardwalk sites (Endeavour Segment) it can be assumed that microorganisms of this group are a component of the subseafloor ecosystem and are advected to the surface from zones with higher temperature. From points (i) and (ii) above it follows that ANME-1GBa organisms are most probably inactive in the surface zone.

To date, there is no direct evidence of the involvement of microorganisms of ANME-1GBa group in the process of anaerobic methane oxidation, although such evidence does exist for representatives of other groups of ANME-1 cluster (7, 27). However, in another our work (32), we detected two closely related mcrA phylotypes (GenBank: HQ635702 and HQ635704) in the FS625 sample studied also in this work (Easter Island vent). The mcrA gene encodes the \( \alpha \) subunit of methyl-coenzyme \( \mathcal{M} \) reductase (MCR), which catalyzes the methane-forming step in methanogenic archaea (53) and is also involved in oxidation of methane catalyzed by microorganisms of ANME clusters (54). Amino acid sequences deduced from the mcrA genes that we detected in the FS625 sample exhibited 91-92% identity to crystallized and well described MCR of methanotrophic
archaea (GenBank: CBH39484 (54, 55)), and our analysis placed these mcrA genes from sample FS625 together with mcrA phylotypes from Guaymas Basin (14) in the root of ANME-1 cluster (Supplemental Material Fig. A2). In the present work, we also detected in the same sample just two closely related 16S rRNA genes affiliated with ANME-1GBa group, and no other ANME-1 phylotypes. Thus, there are grounds to assign the previously detected mcrA sequences to ANME-1GBa group. Considering the similarity of these mcrA sequences to those of methanotrophic archaea, we regard these data as substantial evidence of the methanotrophic nature of the ANME-1GBa microorganisms. Close relatives of mcrA phylotypes that we assign to the ANME-1GBa group were previously detected only in Guaymas Basin hydrothermal sediments (mcrA-Guaymas (14)).

In recent studies Holler et al. (7) and Wankel et al. (16) determined high rates of AOM between 42˚ and 65˚C in incubation experiments. To quantify the abundance of anaerobic methanotrophs, molecular techniques such as qPCR and CARD-FISH were used. But the primers and probes that were used in these studies excluded ANME-1GBa microorganisms. In future studies ANME-1GBa-targeted primers such as ANME1GBHS-183(F) - ANME1GBHS-841(R) ((26) and Table 1) can be applied to both detection and quantification of the ANME-1GBa microorganisms. Further manipulation of vent samples or samples of Guaymas Basin sediments in incubation experiments over a broad range of temperatures coupled with such molecular tools may provide direct evidence of moderate or extreme thermophily of ANME-1GBa microorganisms.

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References


Anaerobic oxidation of methane at different temperature regimes in Guaymas Basin hydrothermal sediments. ISME J. 6(5):1018-1031.


Subbotina IV, Chernyh NA, Sokolova TG, Kublanov IV, Bonch-Osmolovskaya EA,


47. Johnson HP, Embley RW. 1990 Axial Seamount: An active ridge axis volcano on the
central Juan de Fuca Ridge. J. Geophys. Res. 95(B8):12689-12696.


Figure Legends

**Figure 1.** Correlation between the optimal growth temperatures of archaeal type strains and the G+C contents of their 16S rRNA genes ($P_{GC}$).

**Figure 2.** $P_{GC}$ of different groups of ANME archaebacteria with indication of intervals of standard deviation.

**Figure 3.** Dendrogram showing the phylogenetic position of 16S rRNA gene sequences retrieved from the samples analyzed in this study among selected reference sequences of the domain *Archaea*. Sequences from this study are grouped into 14 98%-similar OTUs and printed in bold type. The tree was constructed in the ARB software package using maximum-likelihood (PHYML) algorithm and non-parametric bootstrap analysis. The percent bootstrap values are based on 100 replicates and are indicated at the nodes with $\geq 50\%$ bootstrap support. Bar=10% estimated sequence divergence.
Table 1. *In silico* specificity test of primers and probes targeting 16S rRNA genes of ANME-1 group and its subgroups

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<th>Oligonucleotide name (orientation) and sequence (5’→3’)</th>
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<th>Results of <em>in silico</em> specificity test</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>ANME-1a, %</td>
</tr>
<tr>
<td>ANME1-305 (F) AGCCCGGAGATGGGTCT</td>
<td>25</td>
<td>96</td>
</tr>
<tr>
<td>ANME1-350 (R) AGTTTTCGCCGCTGATGC</td>
<td></td>
<td>98</td>
</tr>
<tr>
<td>ANME1-632 (F) TCAGGGAATCTGCTTGG</td>
<td></td>
<td>50</td>
</tr>
<tr>
<td>ANME1-830 (R) TCGCAGTAATGCCCAACAC</td>
<td></td>
<td>84</td>
</tr>
<tr>
<td>Mix of primers ANME1-395 (F) AACTCTGAGTGCCCTCCA</td>
<td>26</td>
<td>80</td>
</tr>
<tr>
<td>ANME1-1417 (R) CCTCACCTAAAYCCCACT</td>
<td></td>
<td>63</td>
</tr>
<tr>
<td>ANME1GBHS-183 (F) ATACCTGGAATGGGCGGA</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>ANME1GBHS-841 (R) AACACCGGCAACACTCGT</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>ANME1-1337 (F) AGGTCCTACGGGAGGCAT</td>
<td></td>
<td>94</td>
</tr>
<tr>
<td>ANME1-724 (R) GGTCAAGACGCTTGCCT</td>
<td>27</td>
<td>92</td>
</tr>
<tr>
<td>ANME1-25 (F) GAGGCCYACTGCACTACGAGMT</td>
<td>This study</td>
<td>95</td>
</tr>
<tr>
<td>ANME1-1118 (F) CYCRCAGTTGCGCGATCTC</td>
<td></td>
<td>68</td>
</tr>
<tr>
<td>ANME1-1406 (R) AYCYACCTCGGGTGGCTTGGA</td>
<td></td>
<td>100</td>
</tr>
<tr>
<td>ANME1-1G612 (R) CTGGCCCATCGTTTAC</td>
<td>7</td>
<td>16</td>
</tr>
<tr>
<td>cANME1-1G612 (R) CTAGGCCGACTCAGTTC</td>
<td></td>
<td>56</td>
</tr>
<tr>
<td>ANME1-1G1186 (R) GGACATCCTGCATCCAG</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>ANME1-628 (F) GCTTTCAAGGGAATCTGC</td>
<td>28</td>
<td>38</td>
</tr>
</tbody>
</table>
Notes: The results of in silico specificity tests are presented as percentage of perfectly matching sequences, and, for negative controls, as a series of three numbers, the first of which shows percentage of perfectly matching sequences, and the second and third show the percentage of sequences exhibiting one and two mismatches, respectively. As negative controls, we used the 16S rRNA gene sequences of the type strains of archaeal species available from the RDP database (RDP Release 10, Update 28, Jan 12, 2012), representing 342 out of the 374 currently recognized archaeal species.
Table 2. Description of samples that yielded positive results of ANME-1 16S rRNA gene amplification

<table>
<thead>
<tr>
<th>Sample</th>
<th>Geographical location, site and vent</th>
<th>Coordinates and depth</th>
<th>T, °C</th>
<th>Total cell conc.</th>
<th>Primer set 1</th>
<th>Size of the clonal libraries</th>
<th>Detected ANME-1 groups (% of clone library)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MC118</td>
<td>Gulf of Mexico, Mississippi Canyon 118</td>
<td>28°,51.47’N, 88°,29.52’W, 880 m</td>
<td>5</td>
<td>N.D.</td>
<td>+2</td>
<td>1+1+2</td>
<td>24</td>
</tr>
<tr>
<td>FS611</td>
<td>JFR1, Axial Seamount, Bag City</td>
<td>45°,54.974’N, 129°,59.354’W, 1410 m</td>
<td>11.2</td>
<td>1.7 × 10⁴ ml⁻¹</td>
<td>N.D.</td>
<td>+1</td>
<td>96</td>
</tr>
<tr>
<td>FS625</td>
<td>JFR, Endeavour segment, Easter Island</td>
<td>47°,58.88’N, 129°,5.967’W, 2150 m</td>
<td>17.8</td>
<td>1.3 × 10⁴ ml⁻¹</td>
<td>N.D.</td>
<td>+2</td>
<td>96</td>
</tr>
<tr>
<td>FS725</td>
<td>JFR, Endeavour segment, Boardwalk</td>
<td>47°,58.11’N, 129°,5.24’W, 2150 m</td>
<td>16.4</td>
<td>7.1 × 10⁴ ml⁻¹</td>
<td>N.D.</td>
<td>+2</td>
<td>96</td>
</tr>
<tr>
<td>FS448</td>
<td>Mariana arc, NW Rota-1, Fault Shrimp</td>
<td>14°,36.06’N, 144°,46.62’E, 520 m</td>
<td>25</td>
<td>3.49 × 10³ ml⁻¹</td>
<td>N.D.</td>
<td>+1</td>
<td>96</td>
</tr>
<tr>
<td>BG410</td>
<td>Gulf of California, Guaymas Basin BIG 1</td>
<td>27°,00.37’N, 111°,24.56’W, 2000 m</td>
<td>50-70</td>
<td>N.D.</td>
<td>+4</td>
<td>N.D.</td>
<td>28</td>
</tr>
</tbody>
</table>

1 Juan de Fuca Ridge.
2 Total cell concentrations were previously published (32,33).
3 1=ANME-1-25(F) - ANME-1-1406(R), 2= ANME-1-1118(F) - ANME-1-1406(R), 3= ANME-1-25(F) – ARCH-915(R).
4 Results for the clone library constructed by using ANME-1-25(F) - ANME-1-1406(R) primer pair.
The graph shows the relationship between $P_{GC}$ (mol%) and $T_{opt}$ (°C) for Acidiplasma cupricumulus. The equation $P_{GC} = 0.18T_{opt} + 49.8$ with $R^2 = 0.819$ and $n = 355$ is used to model this relationship. The data points represent different temperature zones for the organism:

- **Mesophiles** (<50°C)
- **Moderate thermophiles** (50-70°C)
- **Extreme thermophiles** (70-80°C)
- **Hyperthermophiles** (>80°C)
**ANME phylogenetic groups**

- ANME-1a
- ANM-1AT
- ANME-1GBa
- ANME-1b
- ANME-2ab
- ANME-2c
- ANME-3

**$P_{Gc}$, mol%**

- ANME-1a: 55.7
- ANM-1AT: 57.3
- ANME-1GBa: 63.5
- ANME-1b: 55.2 ± 0.1
- ANME-2ab: 55.3 ± 0.1
- ANME-2c: 55.5 ± 0.1
- ANME-3: 53.4 ± 0.1