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Novel microbial communities of the Haakon Mosby mud volcano and their role as a methane sink

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Abstract:

Mud volcanism is an important natural source of the greenhouse gas methane to hydrosphere and atmosphere^{1,2}. Recent investigations show that the number of active submarine mud volcanoes may be much higher than anticipated (eg. 3-5), and that gas emitted from deep-sea seeps may reach the upper mixed ocean⁶⁻⁸. Unfortunately, global methane emission from active submarine mud volcanoes cannot be quantified because their number and gas release is unknown⁹. Another uncertainty is the efficiency of methane oxidizing microorganisms in methane removal. Here we investigated the methane-emitting Haakon Mosby Mud Volcano (HMMV, Barents Sea, 72°N, 14°44'E; 1250 m water depth), to provide quantitative estimates of in situ composition, distribution and activity of methanotrophs in relation to gas emission. The HMMV hosts three key communities; aerobic methanotrophic bacteria (*Methylococcales*), anaerobic methanotrophic archaea (ANME-2) thriving below siboglinid tubeworms, and a novel clade of archaea (ANME-3) associated with bacterial mats. We found that upward flow of sulphate- and oxygen-free mud volcano fluids restricts the availability of these electron acceptors for methane oxidation, and hence the habitat range of methanotrophs. This mechanism limits the capacity of the microbial methane filter at active marine mud volcanoes to <40% of the total flux.

The HMMV (Fig. 1), a circular structure of 1 km diameter and <10 m elevation above the adjacent seafloor, has been studied since the 1990s as a typical example of an active mud volcano⁹. Its formation may have coincided with a submarine landslide during the late Pleistocene, 330–200 ka before present¹⁰. Today, fluids, gas and muds rise from 2-3 km depth through a conduit below the HMMV^{11,10}. The emitted gas is of a mixed microbial/thermogenic origin and consists of >99% CH4 with a δ^{13} C-isotope signature of -60‰^{12,13}. The rising fluids are depleted in sulphate, chloride and magnesium caused by subsurface clay dewatering¹¹. Investigation of the HMMV with RV POLARSTERN and ROV VICTOR 6000 in 2003 showed extensive outcroppings

of fresh subsurface muds associated with steep thermal gradients¹⁴, gas and fluid vents, and a large gas plume reaching the mixed upper water column above the HMMV^{12,8}. Seafloor videography in combination with geochemical measurements provided *in situ* estimates of gas flux⁸, fluid flow¹⁵ and habitat distribution¹⁶. We focused on the three main concentric habitats above the gassy muds (Fig. 2): the centre of HMMV, which was devoid of epifauna; thiotrophic bacterial mats dominated by a *Beggiatoa* species; and surrounding fields of siboglinid tubeworms. Gas concentrations in sediments and bottom water were elevated in all three habitats (Tab. 1). An essential difference between the habitats are the fluid flow velocities modelled from *in situ* gradients of temperature, oxygen and sulphide concentrations¹⁵. The results suggest that the upward transport of oxidant-depleted mud volcano fluids hinders downward diffusion of oxygen and sulphate, and limits methanotrophic habitats in the centre and below bacterial mats to the uppermost milli- to centimetres, respectively¹⁵. We have now identified and quantified the methanotrophs populating the three habitats to assess their role in controlling methane flux at HMMV (Tab. 1; Figs. 2, 3).

The surface of the most recent mud flows in the centre of the HMMV, characterized by steep temperature gradients of 3°C m⁻¹ (14), hosted high microbial cell numbers, reaching $5.1 \pm 0.3 \times 10^9$ cells cm⁻³, of which 50 (± 5) % belonged to type I aerobic methanotrophic bacteria of the order *Methylococcales* and *Methylophaga* (Figs. 2-3, 3a3). The 16S rRNA gene sequences of the dominant *Methylococcales* types (HMMV-MetI and -MetII) are closely related to sequences of methanotrophic mussel symbionts (94-97%)¹⁷ and to clone sequence Hyd24-1 (96-99%) from gas hydrate bearing sediments of Hydrate Ridge¹⁸. High concentrations of a ¹³C-depleted (-80‰), type I methanotroph specific fatty acid (FA; C16:1 ω 8c) were found in the same samples (Fig. 3a4). As predicted by the fluid flow model for this area¹⁵, methanotroph cell numbers, lipid biomarkers and aerobic methane oxidation rates (MOx) peaked in the uppermost surface sediment and decreased >5-fold in the second cm because of limited

oxygen penetration (Fig. 3a). Only minor amounts of methanotroph lipids (<0.1 μ g g-dw⁻¹) and very low cell numbers (~10⁷ cells cm⁻³) were found below 5 cm sediment depth (Fig. 3a3-3a4). ANME cells were not microscopically detectable in the centre cores using all known ANME oligonucleotide probes, and the anaerobic oxidation of methane (AOM) remained below detection.

In situ microsensor profiles of sulphide concentrations in HMMV sediments suggested that AOM occurs in older mud flows at 2-3 cm below the sulphide-oxidizer mats (Fig. 2-4). Accordingly, microbial biomass peaked in the top 3 cm (Fig. 3b3) and mainly consisted of a novel consortium of archaea and sulphate-reducing bacteria (SRB) forming dense cell aggregates (Fig. 2-4). 16S rRNA gene analyses of DNA revealed a dominant cluster of archaeal sequences forming a new phylogenetic group named ANME-3 (Supplement Fig. 1). Single sequences of this novel cluster were only sporadically detected at cold seeps^{19,20}, which are typically dominated by other phylogenetic clades of anaerobic methanotrophs, i.e. ANME-1 and ANME-2 together with their bacterial of partner, sulphate reducers (SRB) the Desulfosarcina/Desulfococcus (DSS) branch^{21,19,20}. We propose that members of the ANME-3 cluster are also capable of AOM, based on the coinciding AOM maximum with the sulphide production zone, the presence of high numbers of ANME-3 aggregates (20 (\pm 0.6) \times 10⁶ aggregates cm⁻³ at 1-2 cm sediment depth) making up 81% of total cell numbers, and highly elevated lipid biomarker concentrations of sn2hydroxyarchaeol, archaeol and specific penta-methyl-icosenes (Fig. 3b). Moreover, stable C-isotope signatures of the archaeal lipid biomarkers of δ^{13} C-values <-98‰ indicate that methane is the main carbon source of ANME-3 cells. Analyses of 16S rRNA genes and lipid biomarkers showed that the bacterial partner associated with ANME-3 is different from all other ANME consortia (see supplementary information). It is closely related to *Desulfobulbus spp.* (DBB) and also takes up methane-derived carbon as indicated by the δ^{13} C-value of -70% of its specific lipid biomarker C17:1 ω 6c (Fig. 3b4). As predicted by the fluid flow model, 95% of all ANME-3/DBB aggregates were found directly below the *Beggiatoa* mats (Fig. 3b-3). Because of reduced fluid flow velocities, sulphate can penetrate the upper 4-6 cm of the sediment column (Fig. 3b2, Tab 1). Furthermore, the re-oxidation of sulphide by the sulphide oxidizer mats replenishes the sulphate pool in these sediments, further contributing to the maintenance of high activity and biomass of the AOM community in this zone.

The next major transition in HMMV habitats occurs about 300-400 m from its geographical centre, where the sulphide-oxidizer mats are replaced by dense tubeworm colonies on the hummocky periphery of the HMMV (Fig. 2-5, Tab 1). Two siboglinid tubeworm species, Oligobrachia haakonmosbiensis and Sclerolinum contortum¹³ populate this area with biomasses of 1-2 kg wet weight m^{-2} (estimated from sieved boxcore samples). O. haakonmosbiensis, the dominant species in the investigated area, is up to 60 cm long and has a diameter of 0.5 mm. In the tubeworm fields, fluid flow rates modelled from temperature profiles (0.4 m yr^{-1}) were lower than those in the centre or bacterial mat area¹⁴. In agreement with the low subsurface temperature gradient in this zone (Fig. 3b1, 3c1)¹⁴, gravity cores from this area contained gas hydrate below the worm-infested sediment layers. Balancing downward diffusion and upward fluid flow, maximum sulphate penetration could reach 12 cm. However, the measured oxygen and sulphate penetration in the tubeworm field (Fig. 2-5; 3c1-3c2; Tab 1) show that the tubeworms substantially increase the transport of electron acceptors into deeper sediment layers. Accordingly, the maximum in AOM was found between the base of the worm tubes and the hydrate layer (60-90 cm bsf), coinciding with a subsurface peak of ANME-2/DSS aggregates (5.5 \times 10⁶ aggregates) and their specific biomarker lipids (Fig. 3c). Integrated *ex situ* methane consumption was higher than in other zones of HMMV (Tab. 1). Microbe-invertebrate symbioses have an advantage over free-living microbial populations because they can engineer their environment to increase access to both electron donors and acceptors by special migratory behaviour, mining and pumping. It remains an interesting question which factors determine the transition between *Beggiatoa* mats and tubeworm fields, and the temporal succession of the thioand methanotrophic communities. Visual observation of the seafloor showed that sulphide oxidizing mats and tubeworms are mutually exclusive, i.e. no worms were found within any bacterial mats. The tubeworms have a clear advantage over bacterial mats because they can reach much deeper zones of sulphide production, and may efficiently compete for sulphide and methane uptake. Probably tubeworm larvae cannot settle on *Beggiatoa* mats due to the high fluid flow and high AOM-based sulphide concentration present in this habitat.

In conclusion, we found that methane oxidation was the main energy source for microbial biomass in all different habitats of HMMV. Populations of aerobic or anaerobic methanotrophs comprised between 50-80% of the total microbial community (centre and Beggiatoa mat habitats). At their maxima in abundance and activity (Fig. 3), both communities show relatively low^{22} cell-specific rates of methane oxidation, of <0.1 fmol CH₄ per cell and day under atmospheric pressure and an ambient temperature of -1°C. The total integrated activity and biomass was lower for the aerobic methanotrophs than for the AOM communities, as they were limited to the surface of the centre muds by high fluid flow (Tab. 1 and supplementary information). ANME-3 and ANME-2 consortia both dominate areas with lower flow and profit from the association with other organisms re-oxidizing sulphide to sulphate. Total methane consumption in the centre, *Beggiatoa* and tubeworm habitats was up to 0.4, 1.8 and 2.9×10^6 mol yr⁻¹, respectively (sum 5.1×10^6 mol yr⁻¹). In 2003, methane emission to the hydrosphere from HMMV reached at least 8 to 35×10^6 mol yr⁻¹ (8) resulting in a total methane flux of 13 to 40×10^6 mol yr⁻¹. Aerobic oxidation of methane played a minor role (1-3%) compared to anaerobic oxidation, which removed up to 37% of the total methane flux. Hence, the efficiency of microbial methane removal is substantially lower than in other methane-fuelled systems²³⁻²⁵. These data confirm that fluid advection is a main habitatstructuring factor at cold seep ecosystems, by regulating availability of electron acceptors and hence distribution and activity of methanotrophs. It is generally assumed that with increasing upward flow velocities, more dissolved methane is transported to the surface, fuelling increasingly active seep communities (e.g. ²⁶). However, geofluids produced from compaction processes and dewatering are electron-acceptor depleted. High flows of methane-laden fluids introduce a negative switch in the fluid flow/methanotroph activity relationship. Furthermore, if such fluids transport heat from below, they may even suppress formation of hydrate, which represent an important although dynamic - sink for methane on earth²⁷. Consequently, as in the case of the HMMV, methane rising with warm, oxidant-depleted fluids may accumulate in muds and escape as free gas to the hydrosphere. Here we have shown that the efficiency of the microbial filter against methane already decreases considerably at fluid flow rates >0.4 m yr⁻¹, causing increased efflux of methane to the hydrosphere and probably even atmosphere⁸. Velocities of decimetres to metres yr⁻¹ are within the average range of fluid flow measured at cold seeps^{28,29}, hence the described inhibition of microbial methane consumption may be widespread at seeps. Further quantitative in situ measurements of fluid flow and microbial methane consumption are necessary to constrain the relevance and control of methane emission from submarine mud volcanoes and other fluid-flow driven geo-bio-systems.

Methods

Sampling

Our studies took place during two cruises with R/V "L'Atalante" (2001) and R/V "Polarstern" (2003) both equipped with ROV "VICTOR 6000. Sediments were collected with ROV pushcores and gravity cores for on board ("*ex situ*") measurements. Sediment cores as well as *in situ* microsensor profiles were obtained along a radial

transect from the centre to the east of the mud volcano (Fig. 1b). Bottom water was sampled with a bottom water sampler and methane concentrations were determined by gas chromatography⁸. A detailed description of the methods described here is provided in the online supplements.

Sulphate reduction and methane oxidation rates

Microbial rates of aerobic and anaerobic methane oxidation (MOx and AOM, respectively) and sulphate reduction (SR) in sediments were determined *ex situ* using 14 CH₄ and 35 SO₄²⁻ tracers^{25,22}.

Lipid analysis

Lipid biomarker from frozen sediment samples and tubeworm tissue were extracted, separated and derivatised as described previously^{30,22}. Single lipid compounds were analysed by gas chromatography, mass spectrometry and isotope ratio mass spectrometry for quantity, identity and stable carbon isotope ratio, respectively³⁰.

Fluorescence in situ hybridisation (FISH)

Cells of *Methylococcales* as well as ANME-3/DBB and ANME-2/DSS aggregates were quantified by FISH with monolabeled oligonucleotide probes according to previously described methods²⁰.

Sulphate and chloride concentrations

Sulphate concentrations were determined by HPLC analysis from the supernatant of 5 ml sediment fixed with 25 ml zinc acetate solution²⁵. Chloride concentrations were determined from pressure filtered pore water by ion chromatography.

Microsensor measurements

Microsensors (20 μ m tip diameter) for oxygen, H₂S and pH were manufactured and used as described previously¹⁵. Profiles were recorded *in situ* with a spatial resolution of 0.025 cm by deploying a profiler unit (equipped with the sensors) with the ROV or a free falling lander system. A detailed description of in situ data and the geochemical models applied here is provided elsewhere¹⁵.

Supplementary Information accompanies the paper on www.nature.com/nature.

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Author Contributions statement H.N. and T.L. contributed equally to this work. T.L., T.N., K.K. and R.A. carried out the 16S-based analyses and microscopy, H.N. and M.E. the lipid biomarker work, A.B. and H.N. the rate measurements, D.B., E.S. and M.S. the microsensor and geochemical measurements, M.K., M.S., A.B. and J.P.F. the geo- and videographical survey and experimental strategy. A.B., H.N. and D.B. wrote the manuscript text, and H.N., T.L. and K.K. the supplements. All authors discussed the results and commented on the manuscript.

The nucleotide sequence data have been deposited in the EMBL, GenBank and DDBJ nucleotide sequence database under accession numbers AJ704650-AJ704653, AJ704631, AM287206-AM287207.

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Figure 1. The Haakon Mosby Mud Volcano. a) Sediment echosounder image (PARASOUND) of a 1.8 km long profile crossing the mud volcano from south to north (pink line in panel b, 14° 43,8 E). High gas content in the centre sediments considerably decreases signal penetration. The stacks of sedimentary layers of the northern and southern rims may correspond to former mudflows. Black and white bars represent 100m horizontal distances. The y-axis (height) is 24-fold exaggerated. b) High-resolution bathymetric map of the HMMV showing sampling stations (abbreviations in supplementary info). The IFREMER software CARAIBES was used to process the microbathymetry data acquired by the multibeam echosounder (Reson SeaBat 8125 and EM2000, georeferenced by Posidonia USBL positioning) mounted on ROV VICTOR.

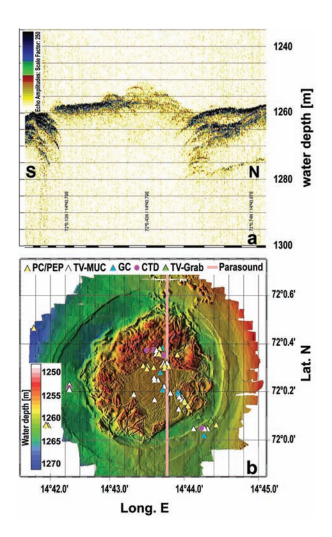
Figure 2. A schematic diagram of the different microbial habitats at HMMV (vertical height is exaggerated 50-fold). (1) Subsurface muds rising from a deep reservoir; (2) Gas and fluid release in the northern centre. The squares (3 to 6) show images of the seafloor (top panel), idealized concentration gradients of oxygen and sulphide (left panel) and micrographs of the dominating microbial organisms (right panel; FISH probe specifications are provided in the online supplements). (3) Central sediments colonized by aerobic methanotrophic bacteria (type I methanotrophs). (4) *Beggiatoa* mats covering sediments dominated by an ANME-3 (red) / DBB (green) pupolation. (5) Colonies of siboglinid tubeworms harbouring symbiotic bacteria (shown in pink; host nuclei shown in blue). (6) Reference station (Micrograph: DAPI staining).

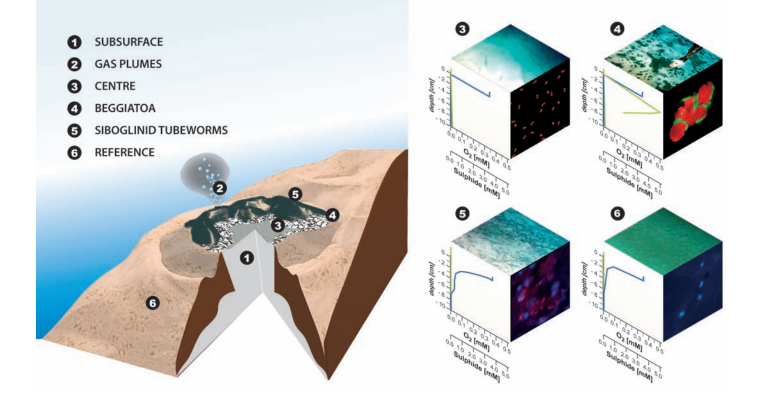
Figure 3. Vertical profiles of oxygen, sulphide and temperature (a1-c1); sulphate, *ex situ* methane and sulphate turnover (a2-c2), bacteria and

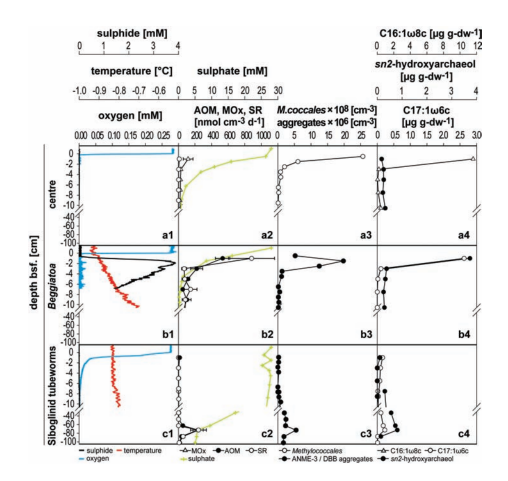
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aggregate counts (a3-c3) and selected lipid biomarker (a4-c4) from the centre (a), *Beggiatoa*-covered sediments (b) and the tubeworm field (c). Oxygen, sulphide and temperature were measured *in situ* with microsensors mounted on a deep-sea profiling unit. AOM, SR, sulphate, cell and aggregate counts as well as lipid biomarker contents were determined using ROV pushcorer samples from the same sites.

Table 1. Gas consumption and emission at HMMV. Total gas emission in 2003 was $8-34 \times 10^6$ mol CH₄ yr⁻¹ based on *in situ* measurements at 3 active vents of HMMV⁸. Maximal relative methane consumption by microbial communities was estimated from microbial methane consumption^{*} divided by the total flux.







| | Centre | Sulphide oxidizer mats | Siboglinid tubeworm fields | Reference |
|--|---------------------------------|---------------------------|--|--------------------------|
| area [km ²] (total = 0.74) ¹⁶ | 0.11 | 0.22 | 0.41 | |
| methane conc. [µM] (20 cm above seafloor, n = 3) | 5.7 (± 1.6) | 0.3 (± 0.1) | 0.7 (± 0.7) | 0.02 (±0.01) |
| fluid flow [m yr ⁻¹] (based on temp. gradients) ^{14,15} | 1.3 – 5.3 | 0.6 – 1 | 0.4 | n.a. |
| fluid flow [m yr ⁻¹] (based on chem. gradients) ¹⁵ | 3 – 6 | 0.3 - 0.6 | n.a. | 0 |
| penetration depth [cm] oxygen (in situ) ¹⁵ | 0.1 – 0.3 | 0.1 – 0.2 | 3 – 10 | 5 |
| penetration depth [cm] sulphate (ex situ) ¹⁵ | 2 | 4 – 6 | 100 | > 15 deeper data n.a. |
| methanotrophs | Methylococcales Methylophaga | ANME-3/DBB | ANME-2/DSS ANME-3/DBB (poss. tubew. symb.) | n.a. |
| methanotrophic biomass [mol C m ⁻²] | 0.04 | 0.6 | 0.4 (80; tubeworm biomass) | n.a. |
| ex situ MOx [mol m ⁻² yr ⁻¹] | 0.9 (± 0.4) | n.a. | 0.2 (± 0.03) | 0.02 (± 0.01) |
| ex situ AOM [mol m ⁻² yr ⁻¹] | 0 | 4.5 (± 1.5) | 7.1 (± 2.0)* | 0.1 (± 0.05) |
| <i>in situ</i> oxygen consumption [mol m ⁻² yr ⁻¹] ¹⁵ | 3.8* | 4.1 | 0.6 (without tubeworm respiration) | 0.3 |
| <i>in situ</i> sulphide production [mol m ⁻² yr ⁻¹] ¹⁵ | 0 | 8.2* | n.a. | 0 |
| areal methane consumption [10 ⁶ mol yr ⁻¹] | 0.4 | 1.8 | 2.9 | |
| rel. methane conumption (% of total flux, 13 - 40 10 ⁶ mol yr ⁻¹) | 1 – 3 | 5 - 14 | 7 – 23 | |

* values used for calculations of areal and relative methane consumption

Supplementary Information

Microbial methane consumption at HMMV

Estimates of total methane flux in the ocean floor range from ca 5×10^{12} mol yr⁻¹ (1) to 30×10^{12} mol yr⁻¹ (assuming a 50% contribution of shelf seas²). Obviously, budgeting of marine methane sources and sinks is still problematic, largely because the leakage of methane from gas hydrates and other major hydrocarbon reservoirs is not known, and the efficiency of the microbial barrier at methane seeps has not been constrained. It was estimated that the anaerobic oxidation of methane flux from the seafloor^{3,1}, whereas the role of aerobic methanotrophic bacteria is minor due to limited oxygen penetration. Here, we have studied an active mud volcano, to obtain a quantitative insight into the distribution and functioning of methanotrophic communities and their role as a methane sink. Only very few other investigations of the microbial ecology and biogeochemistry of active submarine mud volcanoes are available, n⁴⁻⁷, none of which include such a quantitative approach.

Tab. 1 summarizes the biogeochemical characteristics of the three habitats in comparison to the reference site outside of the HMMV (Fig. 2-6). The methane inventory in the bottom waters above the different habitats shows substantially higher methane concentrations above the centre, as a consequence of the reduced filtering capacity of the aerobic methanotrophs inhabiting only the thin surface horizon of the fresh muds. The O₂ influx to this central area was 3.8 mol m⁻² yr⁻¹ as determined by *in situ* microsensor profiling⁸, equivalent to a consumption of up to 1.9 mol CH₄ m⁻² yr⁻¹ (molar MOx stoichiometry 1CH₄ : 2O₂). In comparison, methane consumption below

sulphide oxidizer mats calculated from *in situ* sulphide production was 8.2 mol m⁻² yr⁻¹ (molar AOM stoichiometry 1CH₄ : 1H₂S), i.e. 5 times higher than average aerobic methane oxidation rates in the centre. Integrated *ex situ* MOx and AOM rates confirmed this trend, with 0.9 mol CH₄ m⁻² yr⁻¹ in the centre compared to 4.5 mol CH₄ m⁻² yr⁻¹ in the sulfide-oxidizer mats (Tab. 1). *Ex situ* rates were 2-fold lower than the rates calculated from *in situ* H₂S production and O₂ consumption, probably due to degassing of methane after retrieval and incubation at atmospheric pressure.

16S rRNA gene analyses

For comparative sequence analysis, a total of 67 clones carrying archaeal 16S rRNA gene sequences were obtained from HMMV near surface sediments covered with Beggiatoa mats. All sequences grouped and formed a new cluster within the order Methanosarcinales, termed ANME-3. This cluster has a distinct phylogenetic position and includes sequences from other methane seeps: the Eel River Basin⁹, Hydrate Ridge¹⁰, and Eastern Mediterranean Mud Volcanoes (Heijs et al., database release). All yet known ANME-3 16S rRNA gene sequences are highly similar (95-100%) and targeted by probe ANME3-1249 as indicated by the brackets in Fig. S-1. Closest relatives were obtained from Monterey Canyon seep sediments¹¹ (95-98%), an AOM bioreactor with Monterey Canyon seep sediments¹² (94-97%), and a carbonate chimney of the Lost City hydrothermal vent field¹³ (93-94%). Intriguingly, ANME-3 16S rRNA sequences closely related gene were more to cultivated methanogens (Methanococcoides spp., and Methanolobus spp., 95-96%) than any other ANME group.

Interestingly, ANME-3 form dense aggregates with a bacterial partner (Fig. 2-4), just as their sister groups ANME-1 and -2^{14,9,15,10}, and a novel nitrite-dependent AOM cluster¹⁶. The sulphate-reducing bacteria associated with ANME-3 form a new cluster of uncultivated bacteria within the *Deltaproteobacteria* closely affiliated with *Desulfobulbus* spp. (DBB) (Fig. S-2). In contrast, the bacterial partners of ANME-1 and ANME-2 belong to the *Desulfosarcina/Desulfococcus* clade (DSS)¹⁷. The ANME-3 associated DBB cluster has a distinct phylogenetic position and includes sequences from two other methane seeps: the Guaymas Basin and a Mediterranean Mud Volcano but also sequences from other aquatic sediments, and from electrochemical enrichments.

Lipid biomarker signatures of the ANME-3 consortium

The sediments below *Beggiatoa* mats of HMMV are the first habitat where ANME-3 populations have been found to dominate microbial biomass. Hence, we investigated the specific lipid biomarker fingerprint of ANME-3 in comparison to ANME-1 and -2. The archaeal lipids at this site were dominated by the glycerol diethers *sn2*-hydroxyarchaeol, archaeol and irregular isoprenoidal pentamethylicosenes (PMI) with four and five double bounds, all strongly depleted in ¹³C (δ^{13} C <-98‰ vs. Pee Dee Belemnite). Neither PMIs with higher degrees of saturation, nor crocetane were detected, all of which are characteristic for ANME-1 and ANME-2 communities^{18,14,19-21}. Hence, a lipid signature with dominant amounts of *sn2*-hydroxyarchaeol relative to archaeol, the presence of PMI's with five and four double bonds and the absence of other PMI's and crocetane is indicative for ANME-3.

16S rRNA gene analysis in combination with FISH shows that a relative of DBB is associated with ANME-3 at HMMV. Accordingly, high amounts of C17:1 ω 6c were

found, which was previously assigned as specific lipid biomarker to DBB^{22,23}. This biomarker was also among the isotopically most depleted bacterial fatty acids retrieved from this site (δ^{13} C: -70‰). In contrast, ANME-1 and ANME-2 are generally associated with sulphate-reducing bacteria of the DSS group^{14,24,17}. DSS are characterised by dominant amounts of the fatty acids ai-C15:0 and C16:1 ω 5c/cyC17:0 ω 5,6 when associated to ANME-1 or ANME-2, respectively^{25,19,20}.

Supplementary method information

Sampling

Sampling stations of POLARSTERN expedition ARK XIX/3b with ROV VICTOR 6000 to HMMV are shown in Fig. 1b. Sediments were retrieved with ROV push cores (PC), TV guided multiple corer cores (TV-MUC), gravity corer cores (GC) and a TV guided grab sampler (TV-Grab). Water samples were obtained just above the sea floor with ROV PEP bottles (PEP) and from the water column with a CTD rosette (CTD) (data not shown here).

Sulphate reduction and methane oxidation rates

Microbial rates of aerobic and anaerobic methane oxidation (MOx and AOM, respectively) and sulphate reduction (SR) were determined *ex situ* from sediment incubations with ¹⁴CH₄ and ³⁵SO₄²⁻ tracers^{26,21}. Sediments from push cores were incubated using the whole core injection technique²⁶, whereas sediment subsamples

from gravity cores were incubated in butyl rubber sealed glass tubes $(6 \text{ ml})^{21}$. The *ex situ* cores contained ca 1.4 mM CH₄ during incubation. Cores and glass tubes were incubated for 24 h at *in situ* temperature in the dark before the SR and AOM (MOx) reactions were stopped with 20% Zn-acetate or 2.5% NaOH solution²⁶, respectively. MOx and AOM were distinguished according to the presence or absence of oxygen and SR during the tracer incubations. MOx, AOM and SR rate measurements on 3-6 replicate cores per area showed a standard error of 25 to 55% of the average value.

Lipid analysis

Lipid extracts were separated and derivatised into fatty acid methyl esters (FAMES), alcohol trimethylsilyl ethers and hydrocarbons according to previously described methods^{25,21} before injection into a HP 5890 Series II gas chromatograph equipped with a with a 50 m HP5 fused silica capillary column (0.32 mm i.d., 0.17 µm film thickness) and a flame ionisation detector. Chromatographic conditions were as described previously²⁵ with slight modifications of the temperature gradient for the analysis of alcohol trimethylsilyl ethers and hydrocarbons. Column temperature was programmed from 60 to 150 °C at a rate of 10 °C min⁻¹ and then at a rate of 4 °C min⁻¹ to 310 °C (45 min isothermal). Compounds were identified on a Finnigan Trace MS. Stable carbon isotope composition of single compounds were determined using a HP 6890 Series gas chromatograph interfaced via a Combustion Interface III to a Finnigan Delta Plus isotope ratio mass spectrometer. Reported δ^{13} C-values (in per mil deviation from Pee Dee Belemnite, ‰) were corrected for the introduction of additional carbon atoms during derivatisation. δ^{13} C-values have an analytical error of ±1‰.

Fluorescence in situ hybridisation

Novel monolabeled oligonucleotide probes used in this study were MetI-444 (CCTGCCTGTTTTCCTCCC) and MetII-844 (GCTCCACCACTAAGACCT) for HMMV type I methanotrophs, MPH-732 (GTAATGGCCCAGTGAGTC) for Methylophaga related species, and ANME3-1249 (TCGGAGTAGGGACCCATT) for ANME-3 archaea. For Desulfobulbus spp. we used probe 660 (25)and DBB305 (AGTGCCAGTGTGACGGAT). Endosymbionts in O. haakonmosbiensis (Fig. 2-5) were stained with probes EUB338 I-III. Host cells were stained by DAPI.

Sulphate and chloride concentrations

Pore water for sulphate concentration measurements was extracted by centrifugation of 5 ml sediment fixed in corning vials (50 ml) with zinc acetate solution (20%, w/v)²⁶. An aliquot of the supernatant was injected into a Waters HPLC system (Waters 512 HPLC pump, I.C.-Pak anion-column, Waters WAT007355 4.6 x 50 mm, Waters 730 conductivity detector). Isophtalic acid (1mM) was used as a solvent at a constant flow rate of 1ml min⁻¹. Chloride concentrations were determined from pore water extracted by pressure filtration (5 bars) through Teflon squeezers provided with 0.2 µm cellulose acetate filters. Pore water aliquots were analysed using a MetrohmTM 761 Compact IC with chemical suppression, equipped with a 250 × 4 mm ultra high capacity column (MetrosepTM A Supp 5) and a and conductivity detector. A carbonate buffer solvent (3.2 mM Na₂CO₃ / 1 mM NaHCO₃) was used at a flow rate of 0.7 ml min⁻¹. Total sulphate and chloride concentrations were corrected for porosity, which was determined according to a previously described method²⁶.

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Fig. S-1. Phylogenetic tree showing the affiliations of HMMV 16S rRNA gene sequences to selected reference sequences of the domain *Archaea*. Using the ARB software package²⁷ the tree was calculated with nearly full-length sequences by maximum-likelihood analysis in combination with filters, which consider only 50% conserved regions of the 16S rRNA gene of *Archaea*. Branching orders that were not supported by all calculation methods are shown as multifurcations. Partial sequences were subsequently inserted into the reconstructed consensus tree by parsimony criteria, without allowing changes in the overall tree topology. Clone sequences from *Beggiatoa* covered HMMV sediments are in boldface type.

Figure S-2. Phylogenetic tree showing the affiliations of Haakon Mosby Mud Volcano 16S rRNA gene sequences to selected sequences of the *Deltaproteobacteria*. The tree was calculated with nearly full-length sequences by neighbour-joining analysis in combination with filters, which consider only 50% conserved regions of the 16S rRNA gene of *Deltaproteobacteria*. Clone sequences from HMMV sediments are in boldface type.

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