

Correlating microbial community profiles with geochemical data in highly stratified sediments from the Arctic Mid-Ocean Ridge

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Edited by David M. Karl, University of Hawaii, Honolulu, HI, and approved September 5, 2012 (received for review May 4, 2012)

Microbial communities and their associated metabolic activity in marine sediments have a profound impact on global biogeochemical cycles. Their composition and structure are attributed to geochemical and physical factors, but finding direct correlations has remained a challenge. Here we show a significant statistical relationship between variation in geochemical composition and prokaryotic community structure within deep-sea sediments. We obtained comprehensive geochemical data from two gravity cores near the hydrothermal vent field Loki's Castle at the Arctic Mid-Ocean Ridge, in the Norwegian-Greenland Sea. Geochemical properties in the rift valley sediments exhibited strong centimeter-scale stratigraphic variability. Microbial populations were profiled by pyrosequencing from 15 sediment horizons (59,364 16S rRNA gene tags), quantitatively assessed by qPCR, and phylogenetically analyzed. Although the same taxa were generally present in all samples, their relative abundances varied substantially among horizons and fluctuated between Bacteria- and Archaea-dominated communities. By independently summarizing covariance structures of the relative abundance data and geochemical data, using principal components analysis, we found a significant correlation between changes in geochemical composition and changes in community structure. Differences in organic carbon and mineralogy shaped the relative abundance of microbial taxa. We used correlations to build hypotheses about energy metabolisms, particularly of the Deep Sea Archaeal Group, specific Deltaproteobacteria, and sediment lineages of potentially anaerobic Marine Group I Archaea. We demonstrate that total prokaryotic community structure can be directly correlated to geochemistry within these sediments, thus enhancing our understanding of biogeochemical cycling and our ability to predict metabolisms of uncultured microbes in deep-sea sediments.

taxonomic profiling | ultraslow-spreading ridge | amplicon sequencing

Marine sediments host the largest reservoir of organic carbon in the world and outnumber any other environment with respect to microbial cell abundance (1, 2). The microbial activity in this habitat has a profound impact on the global carbon cycle through the remineralization of sedimentary organic carbon, thus ultimately regulating the oxygen level of the atmosphere (3). Extensive sampling and drilling efforts during the past decade have changed our perception of microbial life in the inaccessible deep seafloor fundamentally (4–17). This pioneering work has led to novel insights into the composition and abundance of potentially active groups of Archaea and Bacteria in deep-sea sediments.

The density of prokaryotic cells in coastal and continental margin sediments typically is 10^8 – 10^9 cells/cm³ in the top sediment layers and declines with increasing depth in a logarithmic fashion (2). Even though cell abundance in organic-poor open-ocean sites can be several orders of magnitude lower, it may still exceed

10^5 cells/cm³ at depths close to 1,000 m below the seafloor (mbsf) (12). This omnipresence of prokaryotic cells was demonstrated recently by Roussel et al. (16) who detected viable cells at a sediment depth of 1,626 mbsf, supporting the idea that temperature is the ultimate limit for microbial survival at depth (18). In general, marine sediment communities appear to be dominated by a restricted number of bacterial and archaeal phyla, including Chloroflexi, Planctomycetes, Japanese Sea division 1 (JS-1), a diverse spectrum of Proteobacteria, the Deep Sea Archaeal Group (DSAG), Marine Group I (MG-I), the Miscellaneous Crenarchaeotic Group (MCG), and the South African Goldmine Euryarchaeotal Group (SAGMEG) (reviewed in refs. 13, 19, and 20). These groups have been found in a variety of different marine sediments, including coastal and open-ocean sites, based on 16S rDNA clone libraries (6, 9, 21, 22), 16S rRNA clone libraries (15), and metagenomic shotgun sequencing (23, 24). The vast majority of these communities seem to be alive and active, although the fractions of dead and dormant cells vary substantially among locations (15, 25, 26).

Geochemical pore water profiles in marine sediments show distinct redox zones (27), suggesting that each of these zones are shaped by organisms with specific metabolic traits. In agreement with this notion, several studies have demonstrated down-core stratification of microbial community composition (28–31). Furthermore, a few specific microbial groups, such as anaerobic methane oxidizers (ANME) (32, 33) and anaerobic ammonium oxidizers (anammox) (34) have been associated consistently with specific redox zones. In addition, samples obtained from markedly different sedimentary settings (e.g., high or low temperature, presence or absence of methane, high or low carbon load) have been found to show distinct microbial community compositions (7, 35, 36). Nonetheless, an explicit, quantitative correlation

Author contributions: S.L.J., L.Ø., I.H.S., I.H.T., R.B.P., and C.S. designed research; S.L.J., T.B., K.F., R.F., and I.H.T. performed research; S.L.J., B.H., A.L., and I.H.T. analyzed data; and S.L.J., B.H., I.H.T., and C.S. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

Freely available online through the PNAS open access option.

Data deposition: 16S rRNA gene sequences are deposited in the National Center for Biotechnology Information Sequence Read Archive (accession no. SRP009131). A comma-separated text file listing the taxonomic affiliations of all operational taxonomic units and their distribution across datasets can be downloaded from <http://services.cbu.uib.no/supplementary/jorgensen2012>. Data have been deposited in the Pangaea database (doi:10.1594/PANGAEA.786687).

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1207574109/-DCSupplemental.

between stratigraphic variability in geochemical properties and concomitant changes in the structure of the total microbial community or the relative abundance of individual taxa has remained elusive. This lack of correlation could be explained in part by limited datasets, low spatial resolution, and insufficient depth of the taxonomic profiling. Here we address these challenges and quantitatively explore the relationship between microbial community structure and multivariate geochemistry in highly stratified deep-sea sediments.

We investigated two 3-m-long sediment cores from the ultraslow-spreading Arctic Mid-Ocean Ridge system, one of the few places where substantial amounts of sediment accumulate in the mid-ocean rift valley. The cores were retrieved from two sites, one 15 km southwest (SW) and one 15 km northeast (NE) of the Loki's Castle Vent Field (37). In addition to the hemipelagic and glaciomarine sediments derived from the Bear Island Fan system, the rift valley also receives episodic input of volcanogenic and metallogenic hydrothermal material. The resulting stratification of the cored sediments allowed us to study the influence of changes in sediment geochemistry on the microbial subsurface communities on a compact depth scale. We used deep sequencing of 16S rRNA gene amplicon tags, covering both the archaeal and bacterial domains, to obtain a detailed qualitative and quantitative taxonomic inventory from selected sediment horizons in both cores. We linked estimates of the relative abundance distribution of the entire prokaryotic community to copy numbers of marker genes from both domains as well as to 16 different geochemical and geophysical parameters of sedimentary solids and solutes. We found that stratigraphic variation in the structure of the microbial community as well as in the relative abundance of individual taxa can be correlated directly to stratigraphic variation in geochemical properties across both cores.

Results

Core Descriptions. Core GC6, retrieved SW of the vent field, consisted of highly stratified hemipelagic-glaciomarine sediments with layers of different colors varying on a centimeter scale from light to darker brown and gray (Fig. 1A). Distinct coarser layers of detrital pyrite, likely of hydrothermal origin [71 and 95 cm below sea floor (cmbsf)], and of altered volcanoclastic material (65 and 232 cmbsf) generally were reflected by Fe enrichments in X-ray fluorescence (XRF) core scanner profiles (Fig. 1A) and by elevated Fe₂O₃ and S concentrations in quantitative geochemical sediment analyses (Table 1). The XRF profiles also revealed five distinct Mn-enriched layers, the uppermost of which (22 cmbsf) was interpreted as the lower boundary of the oxic layer (Table 1). Overall, the reduction potential (Eh) profile in GC6 (Table 1) covaries with the color changes, with higher values in the brown sections and lower but still positive values in the gray sections.

Core GC12, retrieved NE of the vent field, consisted of hemipelagic-glaciomarine sediments in the upper half and glaciogenic debris flows in the lower half. The oxic/anoxic transition zone was marked by a single Mn-enriched layer at 38 cmbsf, and the Eh profile again covaried with the sediment color (Table 1).

Total organic carbon (TOC) content varied between 0.1–1.2% of weight in GC6 (Table 1, Fig. 1A) and 0.2–1.3% of weight in GC12 (Table 1 and Fig. 1B). These values were slightly higher than commonly reported for open-ocean sediments and likely reflect input from the Bear Island Fan and the high productivity of the Arctic Ocean. However, these sediments still are considered a low organic carbon system (<1%) compared with continental margin sites.

We recognized four to five chemical redox zones as defined by Canfield and Thamdrup (38) based on pore water depth profiles of dissolved nitrate, ammonium, manganese, iron, and sulfate (Table 1 and Fig. 1). These profiles include an oxic, nitrogenous, manganous, ferruginous, and sulfidic zone (Fig. 1, *Left*), the last deduced from sulfate depletion in GC12. Despite their proximity,

the two cores showed clear differences: Although dissolved Mn²⁺ followed a typical diagenetic profile in both cores, the concentration in GC6 was higher than usually reported for deep-sea sites (up to 200 μM at 156 cmbsf) even in metal-rich deep-sea sediments (39). An increase in NH₄⁺ coincided with the detection of dissolved Mn²⁺ in both cores, likely as a result of oxygen depletion. However, the down-core increase was less pronounced for GC6 and even decreased in the two volcanoclastic horizons. In GC6 dissolved Fe²⁺ increased in the deepest horizon as Mn²⁺ decreased, probably defining the shift from the manganous to the ferruginous zone. In GC12, on the other hand, Fe²⁺ already was elevated in the manganous zone and increased with depth in an irregular manner. A decrease in SO₄²⁻ to 23 mM in the lower part of GC12 indicates microbial sulfate reduction. In contrast, no indication of sulfate reduction was observed in GC6. However, traces of H₂S were detected at 88 cmbsf, between the two detrital pyrite layers. This horizon also showed traces of dissolved Fe²⁺ and a significant pH increase and NH₄⁺ decrease.

NO₃⁻ was depleted from the interstitial phase below the oxic layers. Interestingly, however, NO₃⁻ could be extracted from the solid phase of all horizons in both cores (Table 1), suggesting that it was adsorbed to mineral phases, as observed in soils (40).

Composition and Diversity of the Microbial Community. For each of the 15 sampled sediment horizons, we generated a 16S rRNA gene amplicon library, with one primer set covering the V5–V8 region of both bacterial and archaeal taxa (*Materials and Methods*). Pyrosequencing yielded a total of 59,364 high-quality sequence reads after extensive filtering. In total, 4,790 reads were unique, with an average length of 231 bp.

From the nine horizons in core GC6, 1,668 different operational taxonomic units (OTUs) (cutoff, 97% sequence identity) were divided into 50 classes (39 bacterial and 11 archaeal) representing 33 different phyla. For the six horizons in core GC12, 1,135 OTUs were identified within 48 different classes (36 bacterial and 12 archaeal) distributed among 38 phyla (for a complete list of represented taxa and their abundance on phylum and class level, see Table S1). A comma-separated text file listing the taxonomic affiliations of all OTUs and their distribution across datasets can be downloaded from <http://services.cbu.uib.no/supplementary/jorgensen2012>. At the class rank, more than 85% of all observed taxa were present in both cores, but their abundance varied greatly between cores and among horizons within each core (Fig. 1 and Table S1). Approximately 50% of the OTUs present in core GC12 were shared with GC6. It is noteworthy that none of the 2,277 unique OTUs was represented in all 15 horizons.

Determination of Bacterial and Archaeal Abundance. Quantitative PCR (qPCR) was used to estimate total numbers of bacterial and archaeal small subunit (SSU) rRNA genes separately (Table 1). The total numbers in GC12 (Archaea plus Bacteria) varied between 5.1×10^5 and 5.2×10^6 16S rRNA gene copies/g sediment (wet weight), comparable to the abundance in other open-ocean sediments (41, 42). GC6, on the other hand, exhibited stronger variability between horizons, with both Bacteria and Archaea up to 100-fold more abundant in the horizon where H₂S and Fe²⁺ were detected in the pore water (3×10^8 16S rRNA gene copies/g sediment in total at 88 cmbsf). In contrast, abundances were barely above the detection limit in the volcanoclastic layers.

The archaeal 16S rRNA gene copy numbers made up 3–89% of the total number of SSU rRNA gene copies in core GC6 and 31–75% in core GC12 (Fig. S1). Thus, the microbial population fluctuates between a bacterial- and an archaeal-dominated community even within these relatively short cores. The ratio between bacterial and archaeal SSU rRNA gene copies as determined by qPCR supported the relative amount of Bacteria versus Archaea as estimated from the amplicon library (Fig. S1), indicating that

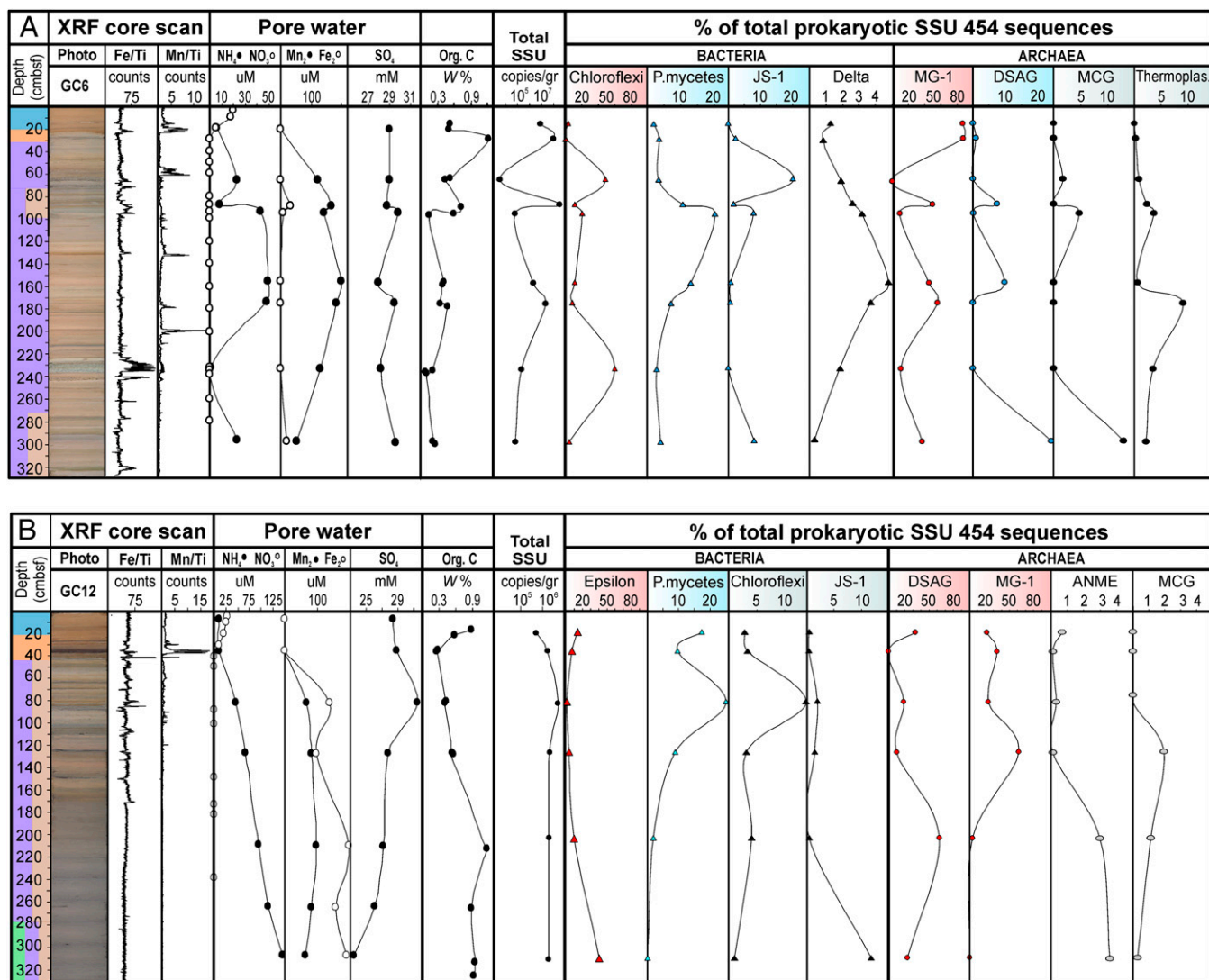


Fig. 1. Characteristics of gravity cores GC6 (A) and GC12 (B) including geochemical data and relative abundances of the four most dominant bacterial and archaeal taxa/phyla. (Left to Right) Photograph of the archive half core; XRF core scanner maps of normalized iron and manganese content; pore water concentrations of ammonium, nitrate, manganese, iron and sulfate; organic carbon content in the sediment (weight %); total number of 16S rRNA gene copies/g sediment (wet weight) as measured by qPCR; percent of total SSU reads obtained from the given taxa in the amplicon library in each horizon. Note that different scales on the x-axis are color coded to indicate the different respiration processes, based on pore water geochemistry: blue, aerobic oxidation; red, nitrate reduction; purple, manganese reduction; brown, iron reduction; green, sulfate reduction. Delta, Deltaproteobacteria; Epsilon, Epsilonproteobacteria; P.mycetes, Planctomycetes; Thermoplas, Thermoplasmata.

the primers used for the amplification for deep sequencing allowed a high rate of coverage and amplified with relatively little bias under the applied conditions.

Principal Components and Correlation Analyses. Principal component analysis (PCA) was performed independently on the relative abundance data (Table S1) and on the geochemical data excluding all gene copy numbers, but including depth (Table 1). Sample scores on the first principal component (PC1) of the two datasets showed a significant rank-order correlation at the class level ($\rho = 0.671$, $P = 0.02$) (Fig. S2A), and the strength of this relationship increased substantially when oxic layers were removed ($\rho = -0.883$, $P = 0.003$) (Fig. S2B). This result demonstrates that changes in the geochemical structure of the sediments, especially below the oxic zone (PC1 explained 41% of the variance), covary with changes in the overall community structure (PC1 explained 33% of the variance at class level).

Next, we plotted the original data for each individual geochemical parameter against PC1 scores of the relative abundance data and found the strongest correlations with TOC ($\rho = 0.621$, $P = 0.016$) (Fig. 2A), dissolved SO_4^{2-} in the pore water ($\rho = 0.691$, $P = 0.006$) (Fig. 2B), and the relative content of iron and manganese in the solid phase [$\rho = -0.821$, $P < 0.000$ (Fig. 2C) and $\rho = 0.582$, $P = 0.025$ (Fig. 2D), respectively].

Finally, pairwise comparisons uncovered several strong linkages between the relative abundance of individual taxonomic groups and specific geochemical parameters (Fig. S3) as well as striking patterns of co-occurrence among microbial groups at different taxonomic levels (Fig. S2 C–F). The most important results are discussed in more detail below.

Distribution of the Most Abundant Bacterial Phyla. Expressing the relative abundance of taxa as the proportion of total SSU rRNA gene tags allows the depth variation of the most abundant bac-

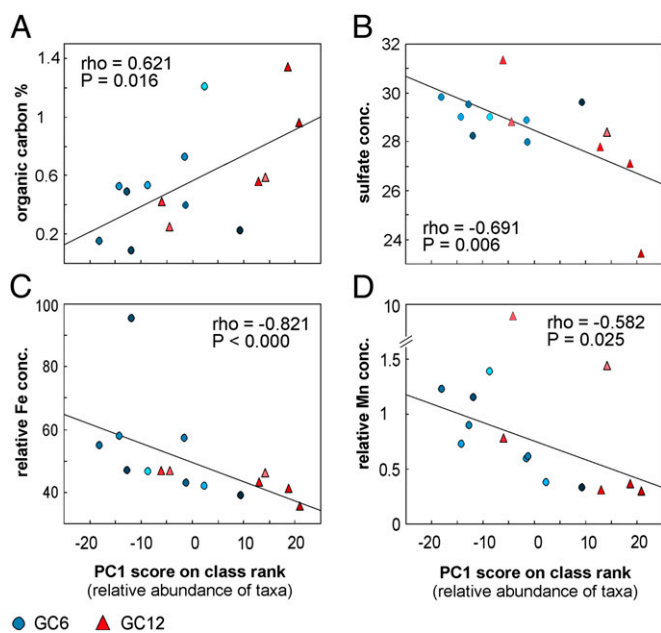


Fig. 2. Significant correlations ($\alpha = 0.05$) between variation in microbial community structure and context data. Microbial community variation is measured by PC1 scores on relative abundance data at the class level. (A) Organic carbon content (% C). (B) Pore water sulfate concentration (mM). (C) Relative iron content in solid-phase iron values as measured by counts by XRF and normalized to Ti counts. (D) Relative content of manganese measured in the solid phase. Manganese values are measured as counts by XRF and normalized to titanium (Ti) counts. Correlations are given as Spearman's rank-order correlation (ρ). Blue circles indicate values from gravity core GC6; red triangles indicate values from gravity core GC12. Color shading indicates depth in sediment (light, shallow; dark, deep).

Distribution of the Most Abundant Archaeal Phyla. As observed for the Bacteria, most archaeal taxa were present (at the class level) in both cores, but the relative abundance varied greatly between and within cores (Fig. 1).

MG-I, also named “Marine Group I.1a,” now assigned to the Thaumarchaeota phylum (46, 47), was the most abundant in six of the nine horizons in GC6. This group contributed as much as $\geq 87\%$ of the total number of SSU rRNA gene-sequence reads in the two top layers (16 and 29 cmbsf). They also dominated in GC12 except for the two deepest horizons (203 and 310 cmbsf). These Archaea not only constitute one of the most abundant microbial planktonic groups in the oceans (48, 49) but also are a highly abundant component in marine sediments (29, 50, 51). Although only aerobic growth has been reported for Thaumarchaeota thus far (52–55), we found MG-I-related organisms in anoxic horizons in our study. Their presence was particularly evident at 126 cmbsf in GC12 (61% of total reads) and at 88, 156, and 174 cmbsf in GC6 (45–56% of total reads), giving an absolute estimate of 4.2×10^5 to 1.5×10^8 MG-I-affiliated 16S rRNA gene copies/g sediment (wet weight) in these horizons. Phylogenetic analysis (Fig. 3) of all published full-length sequences of MG-I for which the habitat origin was assigned unambiguously in the database entry (*Materials and Methods*) demonstrated a separation of planktonic (gamma, delta), sponge-associated (beta), and terrestrial (lambda I, II) taxa from those found predominantly in the sediment (upsilon, eta, iota, theta, epsilon, zeta, and mu clusters; no sequences related to cluster kappa fulfilled the requirements mentioned in *Materials and Methods*) (Fig. 3A). With the exception of sequences affiliated with cluster alpha 1 (9%), all MG-I-related sequences in GC6 and GC12 were affiliated with the lineages that are found predominately in sediments (Fig. 3B and C). This finding augments the earlier

observations by Durbin and Teske (50), who stated that novel clusters distinct from planktonic MG-I organisms can be discovered in marine sediments. A similar habitat-specific clustering has been shown for the archaeal ammonia monoxygenase (*amoA*) (56, 57). To explore the potential of sedimentary MG-I archaea to oxidize ammonia, we quantified the archaeal *amoA* gene, which is used as a genetic marker for this metabolism. Copy numbers ranged from 1.5×10^3 to 2.4×10^6 and from 1.5×10^4 to 2.9×10^6 /g sediment (wet weight) for GC6 and GC12, respectively (Table 1). We found a significant correlation of archaeal *amoA* gene copy numbers with the abundance of group MG-I ($r = 0.929$, $P < 0.000$) (Fig. 4C and Fig. S3E) and total numbers of archaeal 16S rRNA genes ($r = 0.861$, $P = 0.003$) (Figs. 4B and Fig. S3F) in GC6. Both marker genes (*amoA* and 16S rDNA) were found in the same order of magnitude in all samples. More intriguingly, the MG-I abundances also correlated with total NO_3^- concentration ($r = 0.827$, $P = 0.011$) (Fig. 4A and Fig. S3D) and, to a lesser but still significant extent, with TOC content ($r = 0.692$, $P = 0.039$) (Fig. S3C).

The DSAG, also referred to as “Marine benthic group B,” is the most abundant single group of organisms in deep-sea sediments, along with the MCG (20). They were a dominant archaeal constituent in both cores. The relative abundance of DSAG in GC12 ($\sim 25\%$ of total reads) correlated significantly with both TOC and Fe_2O_3 concentration ($r = 0.869$, $P = 0.025$, and $r = 0.819$, $P = 0.046$, respectively) (Fig. 5A and B and Fig. S3I and J). This observation is consistent with a heterotrophic lifestyle (14) possibly coupled to the reduction of iron oxides rather than to sulfur compounds, as previously proposed (7).

SSU rDNA signatures of the MCG often are found in high numbers in clone libraries from deep-sea sediments [reviewed by Teske and Sørensen (19)]. In our samples they constitute up to $\sim 10\%$ of all prokaryotic SSU rRNA genes in certain layers. This group, like the DSAG, has been proposed to be anaerobic heterotrophs (14) and has no cultured representatives. Unlike the DSAG, however, they populate a wide variety of habitats and exhibit larger diversity at the 16S rDNA level. No correlation for this group with any of the measured geochemical parameters was found within our dataset.

Methanomicrobia SSU rDNA signatures were among the four most dominant of the archaeal classes in GC12. The obtained sequences affiliated with ANME groups 1, 2a, 2b, and 2c, all of which are thought to be involved in anaerobic oxidation of methane (AOM) (58). Both their relative and total abundance increased with depth, reaching a maximum of 3.6% of all reads at the deepest horizon (310 cmbsf).

Notably, we found a tight correlation of ANME-1 phylotypes with both Methylococcales and Campylobacterales (Fig. S2D–F), suggesting a common metabolism or a syntrophic partnership.

Discussion

The proximity of the two investigated sediment cores to hydrothermal active sites combined with sediment input from the Bear Island Fan has resulted in a compact redox zonation profile and high concentrations of dissolved metals in both cores. These properties make the sediments interesting model sites to study the variation of subsurface microorganisms in the context of geochemical changes. There were substantial differences between the two cores. For example, the GC12 core is distinguished from GC6 by a clear signature of shallow sulfate reduction and much higher concentrations of dissolved iron (Table 1). Despite such differences, the PCA analysis (Fig. S2A and B) shows that the samples from both cores share a major axis of variation that links changes in the overall composition of the microbial community to changes in the overall geochemical composition. More specifically, the structure of the prokaryotic communities is coupled to variation in the iron and manganese content of the minerals, TOC, and pore water SO_4^{2-} concentration (Fig. 2).

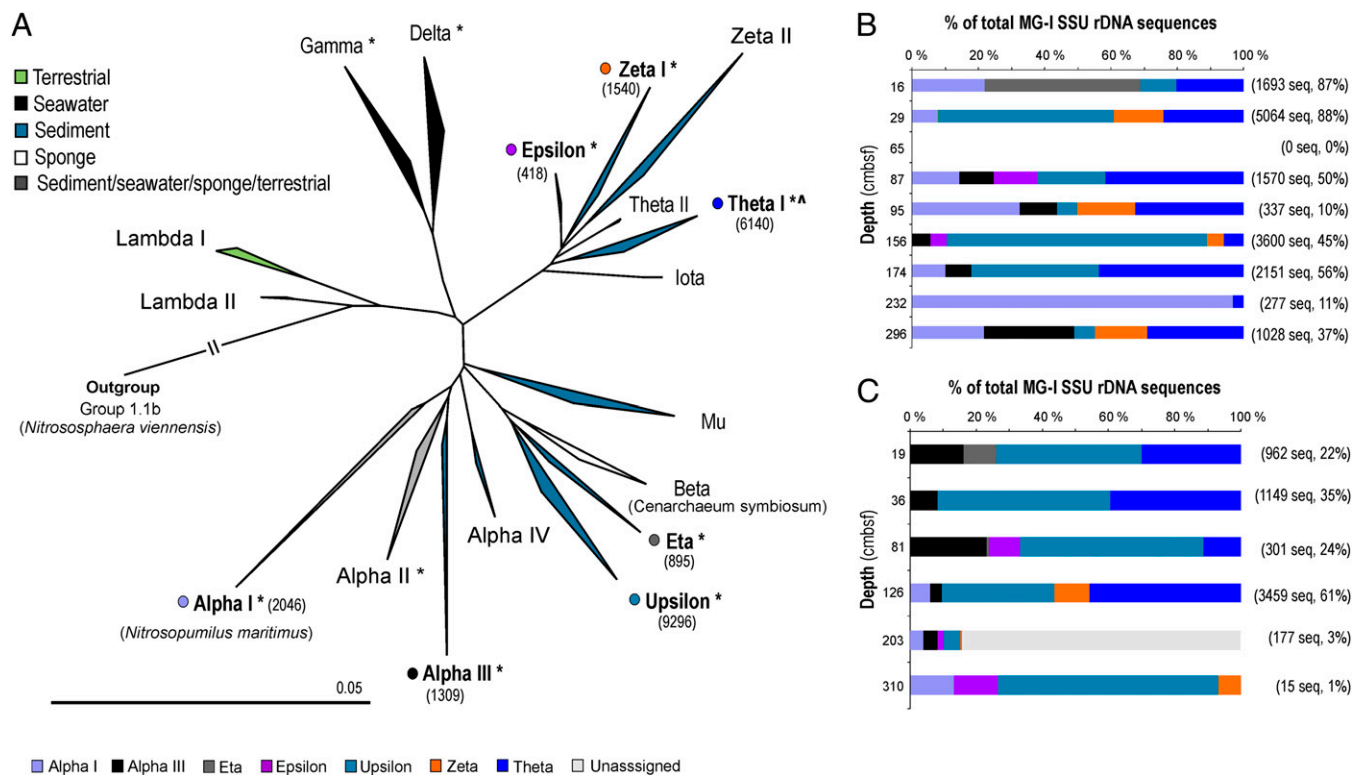


Fig. 3. Phylogenetic analysis and depth distribution of MG-I. (A) Phylogeny based on SSU rRNA gene information from all published sequences available in the Silva database (release 104). The nomenclature follows that used by Durbin and Teske (50), but the additional group names lambda I, lambda II, and mu are given. The tree is reconstructed by NJ using the Felsenstein correction. Topology and clusters are supported by RaxML and PhyML reconstructions on the same dataset. Clusters marked with an asterisk contain sequences retrieved from marine hydrothermal environments. The sediment cluster marked with \cdot contains a subcluster of freshwater/terrestrial sequences. Numbers in parentheses indicate the total number of reads from our study that affiliated with that particular group. (B and C) Depth distribution of MG-1 16S rRNA gene sequences affiliating with each cluster obtained in this study from core GC6 (B) and core GC12 (C). Numbers in parentheses indicate the number of reads from that horizon assigned to MG-1 and the percentage of the total. Color codes correspond to the groups in A.

Our results provide quantitative evidence for the common assumption that organic carbon is one of the fundamental factors shaping microbial communities, supporting the argument that heterotrophic organisms play an important role in deep-sea sediments (14, 20). The content of iron and manganese within the sediments is strongly related to microbial community structure, and, although the causal relationships are highly complex, our results suggest that mineralogy is a key determinant. Finally, we attribute the correlation between abundance and sulfate concentration to the effect of microbial sulfur metabolisms on pore water chemistry, rather than vice versa.

By directly correlating the abundance of each individual taxonomic group to the individual context data, we were able to make predictions about the metabolism of the most dominating organisms in the deep-sea sediments, in particular MG-I, DSAG, and Epsilon- and Deltaproteobacteria. In addition, strong correlations in the relative abundances of different taxa (Fig. S2 C–F) might help elucidate syntrophic partnerships and/or common metabolic preferences.

The archaeal class MG-I belongs to the newly defined phylum Thaumarchaeota, represented by two pure cultures and a few enrichments (59) that were shown to gain energy from aerobic oxidation of NH_4^+ to NO_2^- . Their main carbon source is CO_2 , but coassimilation of organic carbon has been reported for this group (52). The majority of MG-I sequences retrieved in our study (21,644 reads) clustered in phylogenetic groups mainly associated with marine sediments, suggesting the presence of a specialized sediment population (Fig. 3).

None of these groups has any characterized members, but at least four observations support their ability to oxidize ammonia

to nitrite in the sediments from GC6: (i) the tight correlation between the abundance of this group and *amoA* gene copy numbers (Fig. 4C and Fig. S3E); (ii) a high positive correlation of the MG-I with nitrate concentration (Fig. 4A and Fig. S3D); (iii) their co-occurrence with several other phylotypes normally linked to the nitrogen cycle, such as members related to anammox and nitrite-oxidizing *Nitrospina*; and (iv) the fact that the increase in ammonium concentration below the oxic zone is less pronounced than would be expected otherwise (Fig. 1). The presence of an active group of ammonia-oxidizing archaea is surprising, because the sediments are considered to be anoxic below the two uppermost sampling depths. However, MG-I-related sequences have been reported previously from supposedly anoxic environments, although the metabolic implications of these findings have not been addressed (7, 29, 39, 60). Their presence in anoxic horizons can have several explanations: (i) sediment-specific phylogenetic groups of MG-I may have the ability to oxidize ammonium with an alternative electron acceptor; (ii) oxygen could be produced intracellularly, as recently described for the methane-oxidizing group NC10 (50); or (iii) *amo* genes present in the organisms could have some other function, as suggested by Mussmann and colleagues (61). In summary, our data strongly support ammonia oxidation, but the electron acceptor remains unknown. In this context it is noteworthy that members of Methylocoales, another group assumed to be comprised exclusively of aerobes, also occur in highly reduced horizons in this study.

The DSAG is proposed to represent heterotrophic organisms based on indirect evidence from stable carbon isotopes in archaeal lipids (14). A strong correlation between the relative abundance of DSAG and organic carbon content in our study (Fig. 5A and

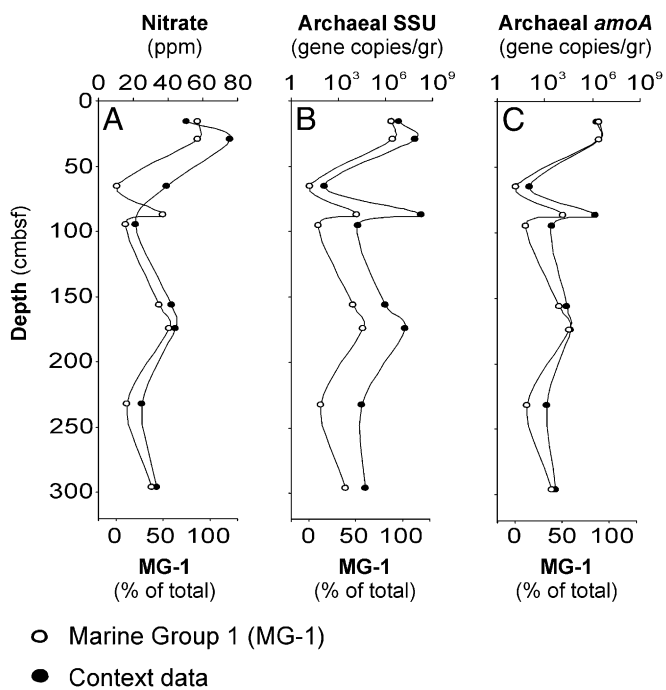


Fig. 4. Covariance between relative abundance of MG-I and context data. The depth distribution of the relative abundance of MG-I in core GC6 strongly covaries with (A) nitrate concentration (ppm) extracted from the solid phase (Pearson's $r = 0.827$, $P = 0.011$); (B) archaeal SSU rDNA ($r = 0.827$, $P = 0.011$); and (C) archaeal *amoA* gene copies ($r = 0.929$, $P < 0.000$). Gene copy numbers are estimated by qPCR and given per gram of sediment (wet weight). See also Fig. S3.

Fig. S3J) supports this suggestion. Their energy-yielding metabolism is a subject of debate, and several studies have proposed direct or indirect coupling to methane oxidation (7, 14, 62) as well as possible sulfate reduction (7). Here we argue that labile iron oxides serve as the terminal electron acceptor either in the direct oxidation of organic carbon or as an energy-yielding metabolism coupled to another electron donor, such as CH_4 , NH_4^+ , or sulfur compounds. We base this proposal on the significant correlation between iron oxide and the relative abundance of DSAG in core GC12 (Fig. S3J). The correlation between DSAG and dissolved Fe^{2+} in the pore water was positive but not significant (Fig. S3K). (GC6 showed a similar relationship but was omitted from statistical analysis because of the low number of observations.)

It is well known that the iron and sulfur cycles are tightly linked; hence Fe^{2+} in the pore water also could result from sulfate reduction to sulfide and the subsequent reaction with iron oxide (63). However, our sulfate profiles do not suggest any significant sulfate reduction except at the deepest horizon in GC12. Thus, iron oxide is the most likely electron acceptor, and the results discussed above, as well as theoretical energy yield, point to organic carbon as the most likely donor. Alternatively, this pathway could involve AOM coupled to iron oxide reduction, a process that potentially yields more energy than AOM coupled to sulfate reduction. Such a metabolism has been demonstrated recently based on geochemical data by Beal and colleagues (64).

In summary, these highly stratified sediments offered a unique possibility to correlate stratigraphic variation in geochemical properties directly to stratigraphic variation in the structure of the microbial community as well as to the relative abundance of individual taxa. We combined deep sequencing of bacterial and archaeal SSU rRNA genes, using a single primer set with broad target coverage, with an extensive set of environmental context data, including nitrate from the solid phase. We used PCA to

distill the covariance structures of the prokaryotic community and the geochemical data separately, thereby revealing a significant correlation between overall changes in geochemistry and overall changes in community composition. Four geochemical components were linked closely to the taxonomic distribution of microorganisms: the total organic carbon, iron, and manganese content in the minerals and the sulfate concentration in pore water. Our findings yield testable predictions about the metabolisms of the most typical and abundant microbial lineages found in the deep subsurface, including the DSAG and sediment lineages of potentially anaerobic MG-1.

Materials and Methods

Site Location and Shipboard Sampling. After retrieval, the cores immediately were cut in sections and split into archive and working halves. Sampling of pore water and sediment for geochemical and microbial analyses from the working halves was conducted onboard the ship as quickly as possible. The two gravity cores, GC6 ~15 km SE of the vent field (73°21.39'N, 7°33.90'E, 3,280 mbsl), and GC12 ~15 km NE of the vent field (73°45.80'N, 8°27.83'E, 3,250 mbsl), were selected for microbiological studies. The in situ bottom water temperature was -0.2°C when cores were retrieved. Samples were collected at depths of 16, 29, 65, 87, 95, 156, 174, 232, and 296 cm in GC6 and 19, 36, 81, 126, 203 and 301 cm in GC12. All samples were collected with sterile 10-mL syringes and were processed immediately or snap-frozen in liquid nitrogen before storage at -80°C . Pore water was extracted with Rhizon samplers from approximately the same depths as the microbiological samples and was analyzed immediately onboard for pH, alkalinity, sulfide, and ammonium or was stored at 4°C until later onshore analysis of dissolved ions. Eh measurements of the sediment were carried out onboard. After sampling was completed, the sediment cores were stored in plastic boxes at $4-6^\circ\text{C}$ on the ship and later at the University of Bergen core repository.

DNA Extraction. DNA was extracted from ~0.5 g of sediment in each sample using a FastDNA spin for soil kit in conjunction with the FastPrep-24 in-

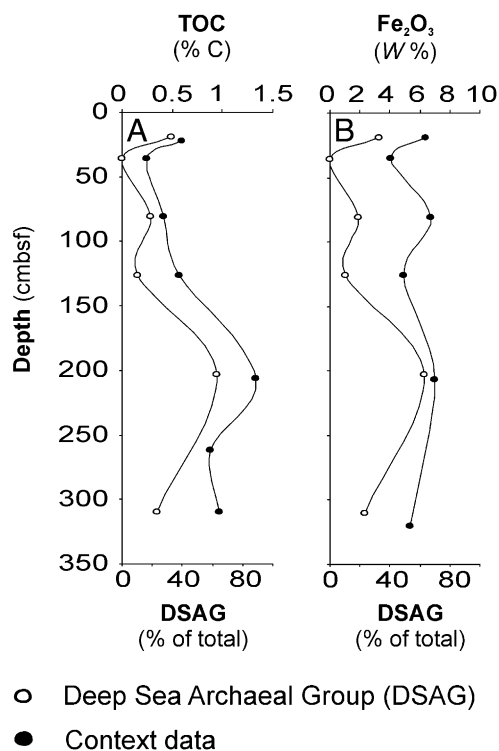


Fig. 5. Covariance between relative abundance of DSAG and context data. The depth distribution of the relative abundance of DSAG in core GC12 strongly covaries with (A) TOC (Pearson's $r = 0.869$, $P = 0.025$) and (B) Fe_2O_3 concentrations in the sediment (Pearson's $r = 0.819$, $P = 0.046$). See also Fig. S3.

strument (MP Biomedicals) following the manufacturer's protocol applying the poly(A) modification described by Hugenholtz et al. (65).

DNA Amplification for the 16S rRNA Gene Library. Different primer combinations were evaluated *in silico* before final selection using the Ribosomal Database Project (RDP) (66), to select the lowest possible degeneracy while maintaining high overall prokaryotic target coverage. In addition, the 454 GS (454 Life Sciences, Roche) FLX technology requires that the length of the amplified product be within 300–800 bp. The optimal primer combination was found to be Uni787F (5'-ATTAGATACCCNGGTAG-3') (67) and Uni1391R (5'-ACGGCGGTGWGTRC-3'), modified from ref. 68. These primers target the V5–V8 region on the 16S rRNA gene and cover 87% and 94%, respectively, of all prokaryotes in the RDP without mismatch (as of May, 2011). The stringency of the performed PCR was kept at a minimum to obtain as much taxonomic diversity as possible, with one mismatch increasing the coverage of both primers to 98%. Furthermore, the chosen primer combination had the least bias toward any one specific taxonomic group ("universal" prokaryotic primers often are problematic, in that they tend to have mismatches against specific taxonomic groups such as Verrucomicrobia, Planctomycetes, and Chloroflexi). DNA from each horizon was PCR amplified in triplicate using the above-mentioned primer combination under the following thermal conditions: 95 °C for 15 min, then 25–30 cycles of 94 °C for 45 s, 53 °C for 45 s, 72 °C for 1 min followed by 72 °C for 7 min before cooling at 4 °C. Each reaction (25 μ L) contained 1 \times HotStar Taq Master Mix (Qiagen), template DNA, and 1.2 μ M of each primer. To ensure correct amplicon length, the PCR product was evaluated by gel electrophoresis. The triplicate PCR products then were pooled to minimize PCR drifting and were purified using GenElute PCR Clean-Up kit (Sigma). A new round of PCR was performed using the same specific primers and thermal conditions but linked to the 454 Life Sciences A and B pyrosequencing adaptor sequence. In addition, the forward primer was labeled with a unique barcode (one for each analyzed sediment horizon) as described by Hamady et al. (69). To minimize PCR bias, the number of cycles was held to a minimum (25–30 in the first PCR and five in the second). Amplicons were purified again, and the concentration and quality were controlled by gel inspection, spectrometry (Cary-300 Bio UV-vis; Varian), and BIO-analyzer (Agilent Technologies). All amplicons (a total of 15 unique samples) then were pooled in a 1:1 ratio based on DNA concentration (~20 ng/ μ L from each sample) and sequenced using multiplex GS FLX pyrosequencing (without titanium chemistry) at the Norwegian High-Throughput Sequencing Centre in Oslo, Norway.

Pyrosequencing flowgrams (SSF files) have been deposited in the National Center for Biotechnology Information (NCBI) Sequence Read Archive under the accession number SRP009131.

Filtering and Removal of Noise from Amplicon Sequence Data. The dataset (84,580 reads) was filtered and cleaned from noise by using AmpliconNoise (70) software. In short, this method includes four steps: filtering, flowgram clustering, sequence clustering, and chimera removal. In the filtering step, reads were truncated at 600 flows, and those with fewer than 360 flows or a noisy signal (flow intensity 0.5–0.7, equivalent to a degenerate base) before this position were removed. In addition, all reads not matching the barcode and primer sequences were removed. In the sequence-clustering step, reads were truncated at 240 bp. AmpliconNoise generated a set of de-noised and chimera-filtered sequences, each with a set of reads most likely to be derived from it. Barcode and primer sequences were removed before further analysis such as taxonomic classification and linkage clustering.

Taxonomic Evaluation. To assign the filtered, de-noised, and chimera-filtered sequences to taxa, we modified and updated the Silva SSUref database release 100 (71) with respect to taxonomy, based on the most recent literature. This database (available at <http://services.cbu.uib.no/supplementary/community-profiling/>) is described in Lanzén et al. (72).

The sequences were aligned to this database using Blastn (NCBI), and the results were analyzed and assigned to specific taxa using the software MEGAN version 3.7 (73) by applying a lowest common ancestor algorithm [for details see Lanzén et al. (72)].

OTU Clustering. All unique de-noised sequences were clustered into OTUs using maximum linkage clustering of pairwise distances. The distance matrix was generated using the Needleman–Wunsch algorithm as implemented in NDist, and clustering was carried out using FDist (both programs are distributed with AmpliconNoise) (70). A 3% distance cutoff was used to define OTUs.

Phylogenetic Analysis of MG-1. The phylogenetic relationship within the MG-1 was evaluated to find habitat-specific groups and to calculate the affiliation

of MG-1 sequences from this study. The tree was calculated based on all published full-length (>900 bp) sequences available in the Silva database release 104 (71), applying the archaeal positional filter and removing highly variable positions (if a sequence was described as "unpublished" in the ARB entry field "journal," it was excluded from the calculation). Length was truncated to ARB position 1773–31131 leaving 784 valid columns for calculation. After calculation, sequences were removed if the ARB entry field "Isolation source" contained one of the following words: mangrove, marsh, or estuary (to avoid uncertainties in the habitat type), leaving 659 sequences. Clusters containing one or more sequences obtained from hydrothermal environments are marked with an asterisk. We do not label these clusters as hydrothermal in Fig. 3, because sequences obtained from marine hydrothermal settings are inherently difficult to assign unambiguously to a specific habitat (e.g., seawater, sediment, chimney, microbial mat). The phylogenetic tree (Fig. 3A) was calculated by Neighbor Joining (NJ) using the Felsenstein correction (74). To verify the tree topology, RaxML (75) and PhyML (76) algorithms likewise were applied on the same dataset; both supported the displayed clustering and grouping. The nomenclature follows that reviewed by Durbin and Teske (50). In addition, we have identified and named three clusters: lambda I and II (exclusively from terrestrial habitats) and mu (exclusively from marine sediments).

All unique sequences from each sediment horizon that could be assigned to MG-1 from our dataset (123 sequences from a total of 21,644) were added to the tree using the parsimony tool in the ARB package. Adding this many short sequences distorts the branch length; hence they are not included in the displayed tree. To confirm the affiliation of our sequences, an NJ tree was calculated based on the above-mentioned database sequences and our sequences but with all truncated to a length of 230 bp. All affiliations were confirmed, with a few exceptions for which the sequences could not be phylogenetically resolved. The depth distributions of our sequences and cluster to which they affiliate are displayed for both gravity core GC6 (Fig. 3B) and gravity core GC12 (Fig. 3C).

Real-Time qPCR. Archaeal 16S rRNA genes were quantified using the prokaryotic primer Uni519F (5'-CAGCMGCCGCGTAA-3') (77) and the archaeal-specific primer Arc908R (5'-CCCGCAATTCCTTAAAGTT-3') [modified from Jurgens et al. (78)]. Each reaction (25 μ L) contained 1 \times QuantiTech Sybr Green PCR master mixture (Qiagen), 0.8 μ M of each primer, and 1 μ L template DNA. The thermal cycling program was 15 min at 95 °C, then 40 cycles of 95 °C for 30 s, 60 °C for 30 s, and 72 °C for 45 s. The quantification standard consisted of a dilution series of a known amount of linearized fosmid 54d9 (79) and a copy number of archaeal 16S rRNA genes between 10 and 10⁷ copies/ μ L [calculated as described by Leininger et al. (80)]. Genomic DNA from *Escherichia coli* was used as a negative control. The R^2 value for the standard curve was 0.99, and the slope value was -3.26 , giving an estimated amplification efficiency of 102%.

Bacterial 16S rRNA genes were quantified using the bacterial-specific primer bac341f (5'-CCTACGGGWWGGCWGCA-3') [modified from Ishii and Fukui (81)] and the prokaryotic 519r (5'-TTACCGGGCKGCTG-3') (77). The quantification standard consisted of a dilution series (between 1 \times 10² and 1 \times 10⁷ copies/ μ L) of a known amount of purified PCR product obtained from genomic *E. coli* DNA by using the bacterial 16S rRNA gene-specific primers 8F/1392R (68, 82). *Sulfolobus solfataricus* genomic DNA was used as negative control. The R^2 value for the standard curve was 0.99, and the slope value was -3.06 , giving an estimated amplification efficiency of 112%. The thermal cycle program was 15 min at 95 °C, then 35 cycles of 95 °C for 15 s, 58 °C for 30 s, and 72 °C for 30 s.

Archaeal *amoA* genes were quantified using the archaeal *amoA*-specific primers CrenamoA23f (5'-ATGGTCTGGCTWAGACG-3') (80) and CrenamoA616r (5'-GCCATCCABCKRTANGTCCA-3') (83). Each reaction (25 μ L) contained 1 \times QuantiTech Sybr Green PCR master mixture (Qiagen), 1.2 μ M of each primer, and 1 μ L template DNA. The thermal cycling program was 15 min at 95 °C, then 40 cycles of 95 °C for 30 s, 50 °C for 45 s, and 72 °C for 45 s. The quantification standard was the same as used in archaeal and crenarchaeal 16S rRNA gene quantification. Genomic DNA from *E. coli* was used as a negative control. The R^2 value for the standard curve was 0.99, and the slope value -3.51 , giving an estimated amplification efficiency of 93%. All qPCR experiments were performed with the Step-OnePlus real-time PCR system (Applied Biosystems) using SYBRGreen as the fluorescent dye. To confirm product specificity, melting curve analyses were performed after each run for all experiments, and each qPCR setup contained samples, standard series, negative controls, and blanks, all in triplicate.

XRF Core Scanning. The archived core halves were scanned using the non-destructive ITRAX XRF core scanner system at Bergen Geoanalytical Facility,

University of Bergen. Samples were irradiated with 3 kW Mo. Step size was 0.5 mm for XRF analysis with a count time of 10 s. Manganese and iron were normalized to titanium counts.

Organic and Inorganic Carbon Measurements. Coulometric titration (CM5012 CO₂ Coulometer; IUC, Inc.) was used to determine the total inorganic carbon (TIC) and the total carbon (TC) contents in 12 of the 15 samples. Analytical precision determined by analysis of replicate standards for TIC and TC was $\pm 0.02\%$ C and $\pm 0.03\%$ C, respectively. The TOC contents were calculated by subtracting TIC contents from the TC contents, leading to an error of $\pm 0.05\%$ C for TOC contents. The remaining three samples were measured applying the method described for nitrogen in the *Solid-Phase Geochemistry* section below.

Pore Water Chemistry. Pore water was extracted using Rhizon samplers, and aliquots were analyzed onboard for pH by using a mobile pH meter, for alkalinity by an autotitrator, and for sulfide and ammonium by spectrophotometric methods (84). Onshore analysis of sulfate was performed by ion chromatography, and dissolved iron and manganese were analyzed by inductively coupled plasma optical emission spectrometry.

Solid-Phase Geochemistry. Eh was measured in all layers directly by electrodes (SP 50x; Consort) inserted into undisturbed sediment as soon as cores were split.

Nitrates were measured in an aqueous solution of (NH₄)₂SO₄ (2 M) and sediment (5:1 ratio) through an ion-selective electrode (nitrate ion combination epoxy electrode no. 31503; Phoenix). Nitrogen and sulfur were determined using an elemental analyzer for CHNS-O based on the principle of dynamic flush combustion coupled with gas chromatography. Nitrogen was measured using a GC packed column for CHNS polytetrafluoroethylene, 2 m (Eurovector), and sulfur was measured using a GC packed column for sulfur, 1 m, 6 × 5 mm (Eurovector). Each sample was dried and powdered before analysis. Iron oxide and manganese oxide contents were determined by atomic absorption spectrometry after digestion of samples by microwave, using H₂O₂+HCl, HNO₃, and HF.

Statistical Analyses. PCA was used to summarize the community structure (relative abundance data) and the geochemical structure (context data) of the sediments quantitatively. Each sample was standardized to zero mean and unit SD. The relative abundance data also were subjected to a centered log-ratio transformation with multiplicative zero replacement (85, 86) to remove any forced correlations imposed by the constant-sum constraint. PCA uses the covariance structure to rotate the original data onto a new set of orthogonal (independent) axes oriented in the direction of maximal remaining variance. By projecting the samples onto the first axis (PC1), we effectively reduced the community (or geochemical) structure to a single variable accounting for the greatest proportion of variance. If the overall community structure were causally related to the overall geochemical structure of the sediment, then we would expect a significant, monotonic (if not strictly linear), arbitrarily signed relationship between the PC1 scores of the relative abundance and geochemical datasets. We tested this hypothesis using the Spearman rank-order correlation (ρ).

The community–geochemistry relationship was dissected further by correlating the relative abundance PC1 scores against the original geochemical measurements, again using Spearman's ρ . Finally, we mined the relative abundance data directly to explore linkages among individual taxa and linkages between individual taxa and geochemical variables, using Pearson's product–moment correlation.

ACKNOWLEDGMENTS. We thank S. Monsen and all the onboard scientific members and crew members of the research vessel *G.O. Sars* during the 2008 H₂Deep expedition. We thank H. Hafliðasson for help with the core scanning and T. Urich for fruitful discussions. The Norwegian High-throughput sequencing Centre at the University of Oslo performed the sequencing (<http://www.sequencing.uio.no>). The project was funded by the H₂Deep project through the European Science Foundation program (EuroMARC). Additional funding was provided by Norwegian Research Council through the Centre for Geobiology.

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