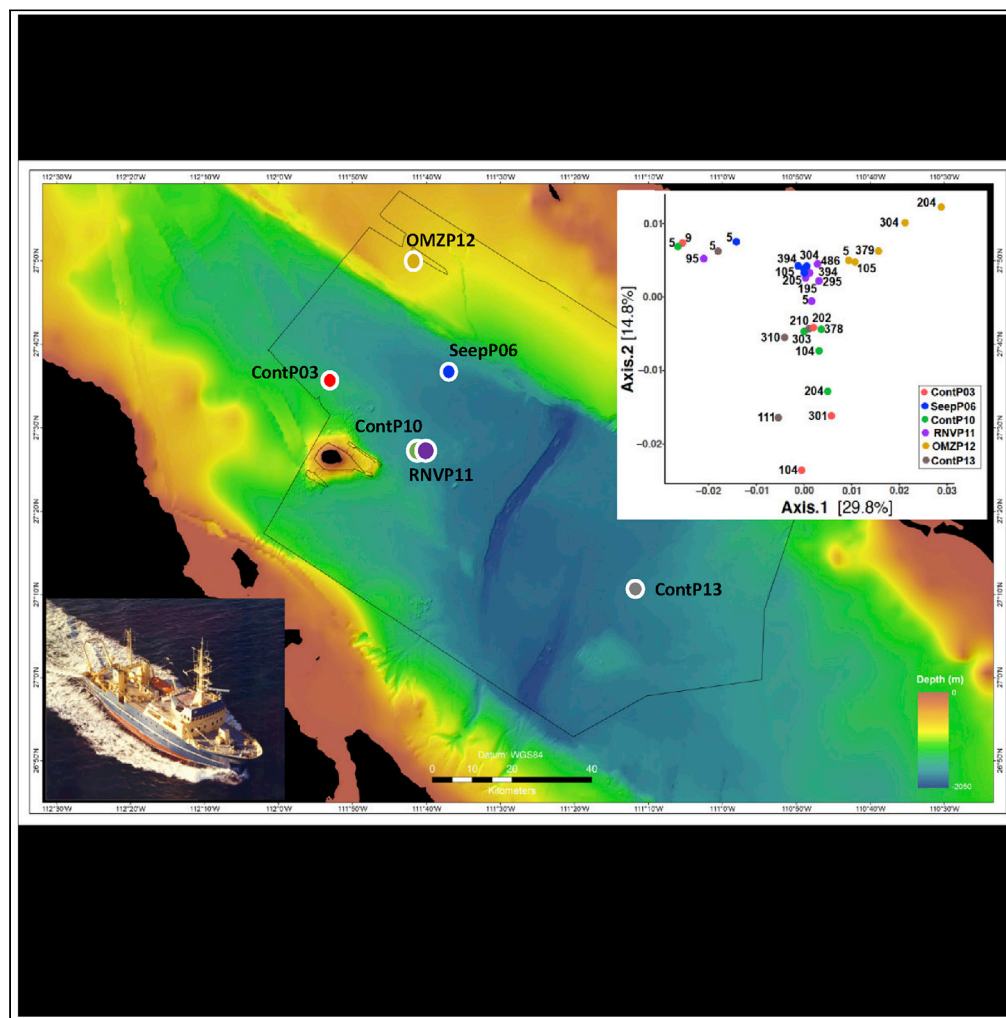


Article

The Guaymas Basin Subseafloor Sedimentary Archaeome Reflects Complex Environmental Histories



Gustavo A. Ramírez, Luke J. McKay, Matthew W. Fields, ..., Christian Hensen, Ana Christina Ravelo, Andreas P. Teske

ramirezg@westernu.edu,
zombiephylogtype@gmail.com

HIGHLIGHTS

Archaeal community composition reflects locally specific environmental challenges

Biogeochemical properties do not predict archaeal community structure

Environmental history controls subseafloor archaeal populations

Ramírez et al., iScience 23, 101459
September 25, 2020 © 2020
The Author(s).
<https://doi.org/10.1016/j.isci.2020.101459>



Article

The Guaymas Basin Subseafloor Sedimentary Archaeome Reflects Complex Environmental Histories

Gustavo A. Ramírez,^{1,2,9,*} Luke J. McKay,^{3,4} Matthew W. Fields,^{4,5} Andrew Buckley,¹ Carlos Mortera,⁶ Christian Hensen,⁷ Ana Christina Ravelo,⁸ and Andreas P. Teske¹

SUMMARY

We explore archaeal distributions in sedimentary subseafloor habitats of Guaymas Basin and the adjacent Sonora Margin, located in the Gulf of California, México. Sampling locations include (1) control sediments without hydrothermal or seep influence, (2) Sonora Margin sediments underlying oxygen minimum zone water, (3) compacted, highly reduced sediments from a pressure ridge with numerous seeps at the base of the Sonora Margin, and (4) sediments impacted by hydrothermal circulation at the off-axis Ringvent site. Generally, archaeal communities largely comprise Bathyarchaeal lineages, members of the Hadesarchaea, MBG-D, TMEG, and ANME-1 groups. Variations in archaeal community composition reflect locally specific environmental challenges. Background sediments are divided into surface and subsurface niches. Overall, the environmental setting and history of a particular site, not isolated biogeochemical properties out of context, control the subseafloor archaeal communities in Guaymas Basin and Sonora Margin sediments.

INTRODUCTION

Guaymas Basin, located in the Gulf of California, México, is a young marginal rift basin where active seafloor spreading generates northeast-to-southwest trending axial troughs surrounded on both sides by extensive flanking regions (Lizarralde et al., 2007). In contrast to mid-ocean spreading centers, axial troughs and flanking regions of Guaymas Basin are covered by thick, organic-rich sediments that represent a combination of terrigenous input and biogenic sedimentation from the highly productive water column (Calvert, 1966). Magmatic intrusions, or sills, are embedded within the thick sediment layers, where they drive hydrothermal circulation (Lonsdale and Becker, 1985) and thermally alter buried organic matter (Seewald et al., 1990), in the process generating complex petroleum compounds (Didyk and Simoneit, 1989), light hydrocarbons and methane (Welhan and Lupton, 1987), carboxylic acids (Martens, 1990), and ammonia (Von Damm et al., 1985). Since the sediments act as a heat-retaining thermal blanket, magmatic activity and organic matter alteration and mobilization are not only limited to the spreading center but also occur at considerable distance, up to 50 km off-axis (Lizarralde et al., 2010). Many of these off-axis sites resemble cold seeps, where methane advection is linked to pathways formed by deeply buried magmatic sills (Geilert et al., 2018). If the underlying sill is sufficiently shallow and hot, the hydrothermal underpinnings of these off-axis sites becomes visible; the recently described Ringvent site provides an example (Teske et al., 2019).

In contrast, the Sonora Margin harbors classical cold seeps where sediment compaction drives reducing, methane-rich seep fluids to the surface. Numerous seep sites with carbonate outcrops and cold seep fauna have been observed on an eroding pressure ridge that follows the transform fault at the base of the Sonora Margin (Simoneit et al., 1990; Paull et al., 2007); the seep communities at these sites are largely based on methanotrophy and sulfide oxidation (Portail et al., 2015). Seep communities and sulfide-oxidizing microbial mats are also widespread on the Sonora Margin slopes (Vigneron et al., 2014; Cruaud et al., 2017).

Finally, most of the extensive flanking regions of Guaymas Basin and the Sonora Margin slope are covered by organic-rich sediments without particular seep or hydrothermal influence; these sediments consist of mixed terrigenous runoff and biogenic components, dominated by diatoms (Calvert, 1966). Sediments on the upper Sonora Margin underlying the oxygen minimum at ca. 600–800 m depth, lack bioturbation

¹Department of Marine Sciences, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA

²College of Veterinary Medicine, Western University of Health Sciences, Pomona, CA, USA

³Department of Land Resources and Environmental Sciences, Montana State University, Bozeman, MT, USA

⁴Center for Biofilm Engineering, Montana State University, Bozeman, MT, USA

⁵Department of Microbiology and Immunology, Montana State University, Bozeman, MT, USA

⁶Instituto de Geofísica, Universidad Nacional Autónoma de México, Coyoacán, México

⁷GEOMAR Helmholtz Centre for Ocean Research, Kiel, Germany

⁸Ocean Sciences Department, University of California at Santa Cruz, Santa Cruz, CA, USA

⁹Lead Contact

*Correspondence: ramirezg@westernu.edu, zombiephylo@gm.com

<https://doi.org/10.1016/j.isci.2020.101459>



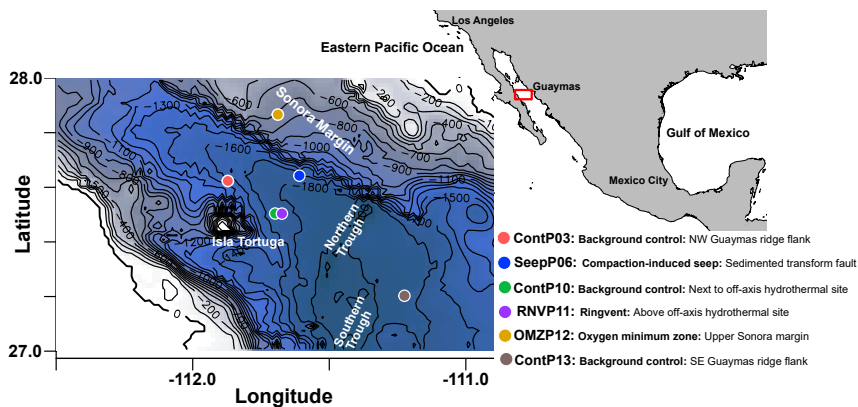


Figure 1. Continental and Bathymetric Hybrid Map Depicting the Location of Guaymas Basin and Sonora Margin in the Gulf of California, and Relevant Coring Sites of the *El Puma* Cruise

The bathymetry blue scale is annotated with 100-m isobaths; the deepest areas in the axial valley range to just below 2,000 m.

and show finely laminated, seasonally changing sedimentation patterns of spring diatom blooms and terrestrial runoff during late summer rains (Calvert, 1964).

Here, to explore links between geochemical and biogeographical patterns in geologically complex settings in the seafloor, we survey the distribution of archaea in diverse sedimentary environments located in the greater Northern Guaymas Basin and Sonora Margin regions. Our samples were collected during a site survey with RV *El Puma* in October 2014 for the recently completed Integrated Ocean Discovery Program Expedition 385 (http://publications.iodp.org/scientific_prospectus/385/index.html). Sampling areas include background sediments from the Guaymas Basin flanking regions, Sonora Margin sediment within the oxygen minimum zone, reducing sediment with cold seep characteristics from the base of the Sonora Margin, and sediment from the off-axis Ringvent site where hydrothermal circulation and methane seepage is driven by a gradually cooling, buried shallow sill. We expand a previous limited sequencing survey of these sediments focused on just one of these sites (Teske et al., 2019) by (1) extending the geochemical analyses, (2) increasing the sampling resolution used for molecular sequencing (from one or two samples per site to ~1-m intervals for all sites), (3) providing a wide breadth of comparative ecological analyses, and (4) discussing the potential implications of our results at a basin-wide scale.

RESULTS

Sediment and Porewater Geochemistry

We surveyed archaeal distribution at six sites on the northwestern and southeastern off-axis regions of Guaymas Basin and on the Upper Sonora Margin (Figure 1, Table S1). These locations represent four different environmental settings: (1) Sediments on the Guaymas Basin flanking regions without hydrothermal or seep activity, represented by cores ContP03, ContP10, and ContP13; (2) the oxygen minimum zone (OMZ) on the upper Sonora Margin (Calvert, 1964), represented by core OMZP12, (3) compacted, highly reducing seep sediments from a pressure ridge, running along the transform fault that is cutting across the base of the Sonora Margin (Paull et al., 2007; Simoneit et al., 1990), represented by core SeepP06, and (4) the Ringvent site, characterized by off-axis hydrothermal circulation (Teske et al., 2019), represented by core RNVP11 (Figure 1). At each site, sediment piston cores ranging from 5 to 486 cm below the seafloor (cmbsf) were collected and geochemically characterized (Figure 2). The sediments are geologically young, ranging in age between ~0.05K and ~20K calendar years, as determined by ^{14}C dating (Teske et al., 2019). The different cores show distinct geochemical characteristics.

Core SeepP06 contains sulfide in millimolar concentrations throughout the core. Below the zone of sulfate-dependent methane oxidation at 1 m depth, methane accumulated to the highest concentrations of this survey, > 10 mM. Porewater Dissolved Inorganic Carbon (DIC) concentrations were consistently high and increased from 15 mM near the interface to nearly 50 mM with depth. These methane and DIC concentrations reached and in part exceeded the highest concentrations previously measured in Sonora Margin seep fluids (Paull et al., 2007). Sediments of core SeepP06 yielded only approximately half of the porewater

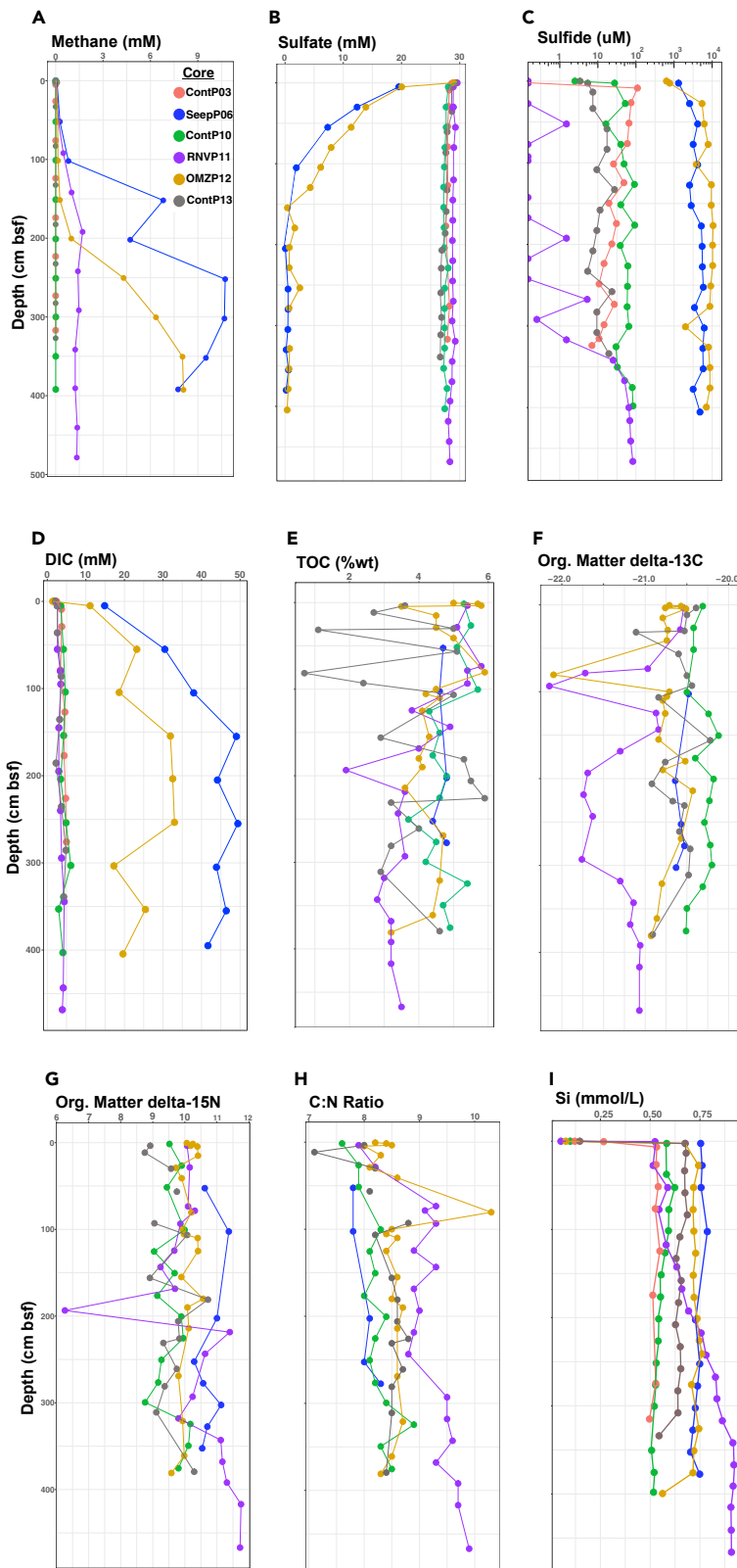


Figure 2. Geochemical Profiles for Guaymas Basin Piston Cores

Geochemical profiles for (A) Methane, (B) Sulfate, (C) Sulfide, and (D) DIC porewater concentrations; (E) Organic Carbon content in weight %; (F) Organic Matter $\delta^{13}\text{C}$ values, and (G) $\delta^{15}\text{N}$ values; (H) Carbon to Nitrogen ratios, and (I) Silica porewater concentrations. Geochemical data for site ContP03 are not available for the analyses depicted in (E)–(H).

volumes of other cores, indicating porewater loss by pressure-induced compaction. Thus, core SeepP06 represents sulfidic, methane- and DIC-soaked, compacted seep sediments from the pressure ridge aligned with the transform fault at the lower Sonora Margin (Simoneit et al., 1990; Paull et al., 2007). Based on diatom assemblages in cemented carbonate rocks, seepage in this area has been ongoing since pre-Holocene times (Paull et al., 2007).

Cores ContP03 and ContP10 share similar methane, sulfate, sulfide, and DIC profiles indicative of non-reducing conditions where sulfate-reducing and methanogenic activities remain minimal and biogenic sulfide and methane occur only in micromolar trace concentrations. With total organic carbon (TOC) between 4 and 6 wt %, the sediments of core ContP10 are organic rich and represent the hemipelagic seafloor sediments of Guaymas Basin that receive ample biogenic sedimentation, mostly by diatoms (Calvert, 1966). The $\delta^{13}\text{C}$ values ranging from -20.12‰ to -20.51‰ are consistent with sedimentary organic material resulting predominantly from phytoplankton input (Teske et al., 2002). Slowly increasing $\delta^{15}\text{N}$ values ranging from 9.04‰ to 10.17‰ and gradually increasing C:N ratios downcore are consistent with microbial utilization of nitrogen compounds in sedimentary biomass.

Contrasting with nearby core ContP10, Core RNVP11 shows the biogeochemical signatures of seawater mixing at Ringvent, such as seawater sulfate concentrations, and previous hydrothermal alteration, as evidenced by silica dissolution and re-precipitation (Teske et al., 2019). Subsurface-derived porewater methane in high concentrations of 1–1.5 mM coexists with porewater sulfate near seawater levels; sulfide is largely absent and reaches 10–100 μM only below 3 m depth. Core RNVP11 also stands out by having the lowest DIC concentrations of all cores, approaching seawater DIC in the upper layers. Below 1 mbsf, organic carbon $\delta^{13}\text{C}$ values are the lowest for all cores ($\delta^{13}\text{C}$ mean of -21.3), whereas C/N ratios are the highest (C/N mean ratio of 9.31), suggesting the influence of isotopically light and nitrogen-depleted hydrocarbons (Peter et al., 1991) (Figure 2). In contrast to other cores, Core RNVP11 shows a strong gradient of dissolved silica, increasing from 0.5–0.6 mM at the surface (similar to ContP10) to 0.9–1 mM at the bottom (Figure 2I). Silica dissolution requires a temperature window of 100°C – 150°C and is therefore considered a marker of hydrothermal activity (Peter and Scott, 1988), ultimately resulting in elevated concentrations of dissolved silica in the water column of Guaymas Basin (Campbell and Gieskes, 1984).

Core OMZP12 differs from all other cores by its location in the oxygen minimum zone on the Sonora Margin slope (Calvert, 1964). Throughout its length, the core showed the conspicuous lamination that is typical for the absence of burrowing infauna and bioturbation in anoxic or severely hypoxic environments. Under these conditions, millimolar concentrations of porewater sulfide permeate the entire sediment core including the surface, otherwise only seen in compaction-induced seep sediments collected at the base of the Sonora Margin (core SeepP06). The sulfate-methane transition zone occurs at approximately 1 and 2 m depth for SeepP06 and OMZP12, respectively. Similar to core SeepP06, porewater DIC increases rapidly with depth, with a maximum value of 33 mM at 254 cmbsf. Sediment TOC, $\delta^{13}\text{C}$, $\delta^{15}\text{N}$, and C:N ratio values generally resemble those of other cores in this survey.

Core ContP13, collected on the southeastern flanking region, differs from other cores by terrestrial input from the Yaqui River. Methane, sulfate, sulfide, and DIC concentrations for this core follow similar depth profiles as observed for cores ContP03 and ContP10. However, TOC varies between ~ 1 and ~ 5 wt % in the first meter of sediment and between ~ 3 and ~ 6 wt % below, suggesting sedimentation pulses of varying organic carbon load. Sediment organic matter $\delta^{13}\text{C}$, $\delta^{15}\text{N}$, and C:N ratios fall within the range of values observed for other cores in this survey.

Diversity of the Guaymas Basin Archaeome

Rarefaction curves are plotted separately for samples in approximately 1-m depth intervals to examine potential downcore trends (Figure 3). Starting at 3 m depth, observed species richness based on rarefaction summaries are lower in Ringvent (Core RNVP11) sediment compared with other sediments (Figure 3, Table S2). Substantially more sequence reads, and thus a higher number of observed species, were recovered

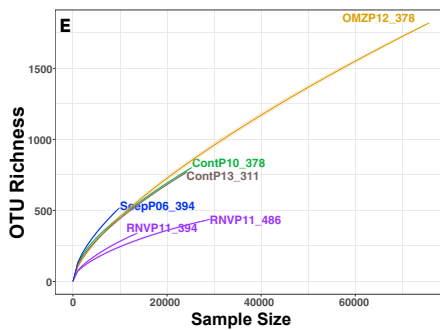
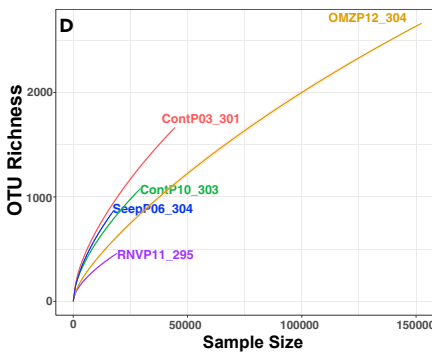
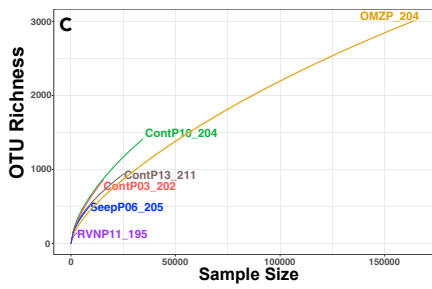
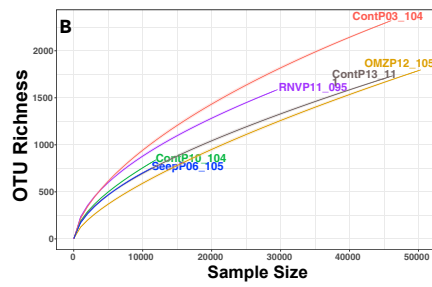
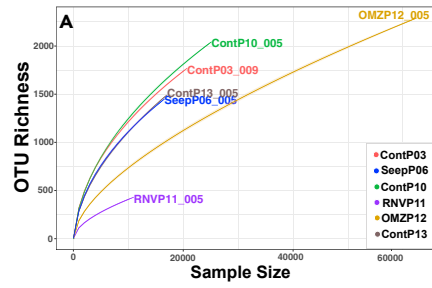


Figure 3. Depth Mapped Rarefaction Summaries (Color Coded to Match Surveyed Sites) for Complete High-Quality Sequence Datasets Depicting Richness as Number of OTUs (97% Similarity Clustered) Observed per Sequences Sampled

(A) Samples near the interface (0–10 cmbsf), (B) samples from ~100 cmbsf, (C) samples from ~200 cmbsf, (D) samples from ~300 cmbsf, and (E) samples from depths greater than or approximately equal 400 cmbsf.

from the Sonora Margin OMZ sediment (Core OMZP12) relative to the other surveyed sites, below 1 m depth (Figures 3C–3E). To account for different sequencing depths without resorting to rarefying the dataset (McMurdie and Holmes, 2014), we estimated total diversity using a non-linear regression model for ratios of consecutive frequency counts, a state-of-the-art method addressing the issue of heterologous sequencing depths affecting richness estimates (Willis and Bunge, 2015). Results from this model indicate no statistically significant ($P_{\text{val}} < 0.05$) differences among surveyed sites (Figure S1).

When beta diversity of the archaeal communities was examined for correlations with environmental metadata using two-dimensional principal coordinate analysis, distinct clustering patterns are observed (Figure 4A). Surface communities for all cores except OMZP12 cluster tightly along negative axis 1 values. All OMZP12 samples cluster along axis 1 values greater than 0.01 independently of sediment depth. Separation along axis 2 partitions the SeepP06, RNVP11, and OMZP12 cores (with positive axis 2 values) from subsurface samples of control cores ContP03, ContP10, and ContP13 with negative axis 2 values; the surface samples of these cores cluster separately (Figure 4A). The influence of environmental factors (i.e., methane, sulfate, sulfide, DIC, water depth, and sediment depth) on community ordination is complex (Figures 4B–4G), and it appears likely that clustering patterns are not driven by these environmental parameters alone. Water column depth (Figure 4F) appears to drive core OMZP12 clustering along larger positive values for axis 1 but most likely represents a proxy for the influence of the OMZ that has persisted throughout the Holocene (Moffitt et al., 2015).

Network Analysis

Network analysis based on the co-occurrence of all operational taxonomic units (OTUs) in each sample reveals that the deepest communities recovered from Core RNVP11 exhibit the greatest degree of separation (Figure 5). At a maximum ecological (Bray-Curtis) distance of 0.8 (i.e., the maximum distance allowed between two samples to be considered connected in the graphical model), most samples share taxa co-occurrence patterns, except for the deepest communities from core RNVP11 (Figure 5A). Decreasing the minimum ecological distance in the model to 0.5 resolves three independent network clusters (Figure 5B). Here, the two deepest samples from core RNVP11 share similar taxa co-occurrence patterns only with each other and are excluded from the two emergent additional networks. In one of these networks, communities near the seawater interface of all cores, with the exception of core RNVP11, connect at no more than 1 degree of separation. Interface sample SeepP06 5cmbsf connects the near-interface sample cluster to all core SeepP06 subseafloor (depth > 1 mbsf) samples. A third independent network shows non-random taxa co-occurrence among subseafloor control sediments (cores ContP03, ContP10, ContP13), a subseafloor and a near-interface sample from core RNVP11, and all samples from core OMZP12; the three deepest OMZP12 samples are only peripherally connected (Figure 5B).

Community Composition

Class-level community descriptions (SILVA132) assigned large membership fractions of the archaeal communities to the Bathyarchaeia, Hadesarchaeaeota, and Thermoplasmata (Figure 6). Asgardarchaea (i.e., sequences classified as Lokiarchaea, specifically) were recovered from every cored site; however, their percent community abundance remained below 0.01% in all samples (Table S2). We note that, when a subset of these samples was studied using a different Archaeal-specific primer set, higher percentages of Lokiarchaea were observed (Teske et al., 2019), implying a potential bias against this lineage in this study. The Class Methanomicrobia, comprising methane-producing and methane-oxidizing members of the Euryarchaeota, was detected in multiple cores but appeared most frequently at depth in core SeepP06. An in-depth summary of the Methanomicrobia reveals the presence of methanogenic families (e.g., Methanomicrobiaceae) and anaerobic, methane-oxidizing ANME lineages (ANME-1, various ANME-2). Notably, ANME-1 archaea dominate core SeepP06 sequence assignments comprising nearly 40% of the total community at 394 cmbsf in this core (Figure S2, Table S3). Order- or higher-level community taxonomic descriptions for all samples generally contained 60% or greater unclassified community fractions (data not shown) when automated taxonomic assignments were performed. In order to not rely on the uncertain output of

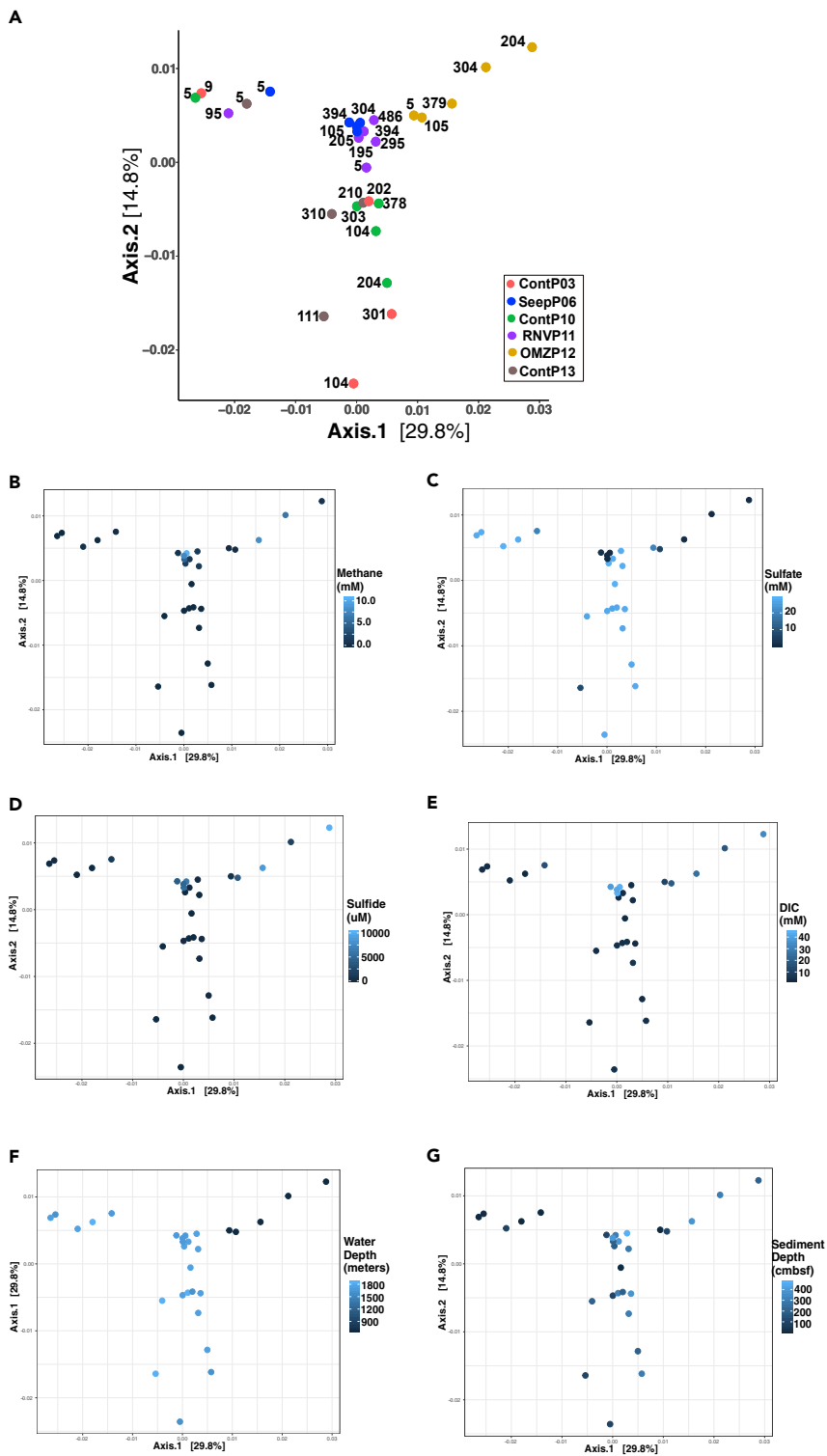


Figure 4. Archaeal Community Grouping Patterns

(A) Two-dimensional Principal Coordinate Analyses of Bray-Curtis dissimilarity distances from r-log normalized sequence count data. Each community plotted is color coded to the core site and numerical labels indicate sediment depth (cmbsf). The first and second axes explain 29.8% and 14.8% of the variance, respectively. Environmental metadata superimposed on ordination plot are (B) methane, (C) sulfate, (D) sulfide, and (E) DIC concentrations, (F) water depth, and (G) sediment depth.

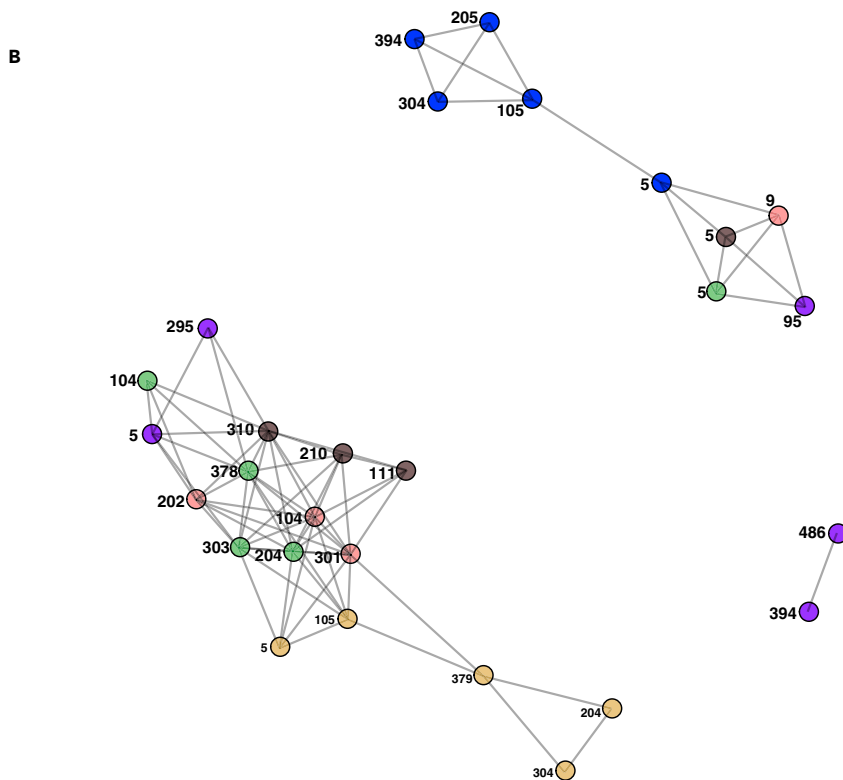
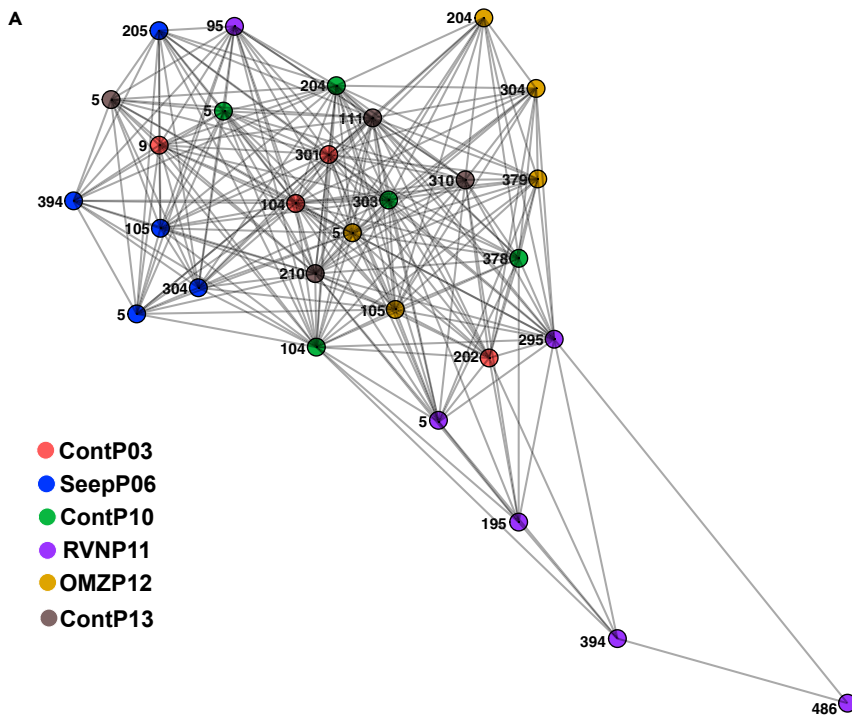


Figure 5. Network Analysis Based on the Co-occurrence of All OTUs at in Each Sample

Nodes represent all archaeal communities analyzed in this study. Nodes are color coded to match descriptions from Figure 1A. Edges are unweighted interactions depicting OTU co-occurrence meeting arbitrary thresholds.

(A) Co-occurrence network threshold set at a maximum Bray-Curtis distance of 0.8.

(B) Co-occurrence network threshold set at a maximum Bray-Curtis distance of 0.5.

taxonomy pipelines, and to resolve archaeal taxonomy assignments in a manner that is consistent with broadly accepted usage (Spang et al., 2017), we also describe community composition based on phylogenetic placement of dominant sequence variants for the most numerous 25 OTUs.

The majority of high-quality sequences in this study (73.0%) clustered into 25 OTU lineages (Figure 7A). Archaeal communities were largely dominated by OTU lineages related to the Bathyarchaea (16 of the top 25 OTUs). OTUs 01 to 03, the three most abundant lineages, belong to the MCG-1, MCG-2, and MCG-3 Bathyarchaea subgroups (Kubo et al., 2012), respectively, with close relatives recovered from Guaymas Basin and globally dispersed seafloor habitats (Figure 7B). High-abundance lineages related to the Marine Benthic Group-D within the Thermoplasmata (MBG-D; OTUs 04, 10, 11, 21, and 23) and the Terrestrial Miscellaneous Euryarchaea Group (TMEG; OTU 08) were recovered from all cores and core depths except the subsurface of Ringvent (Core RNVP11, depth > 1 mbsf). Two highly abundant lineages represented by OTUs 05 and 18 (Figure 7C) were recovered from every core except core SeepP06 and identified as relatives of the Hadesarchaea, formerly known as South African Gold Mine Euryarchaea Group (SAG-MEG)-1 (Baker et al., 2016). Lastly, OTU 14 clustered within anaerobic ANME-1 methanotrophs and was most closely related to ANME-1 phylotypes from cold seep, hydrate, and brine habitats; OTU 14 did not affiliate with the thermophilic ANME-1 Guaymas lineage recovered from hydrothermally active, hot sediments in Guaymas Basin (Biddle et al., 2012) (Figure 7C).

Differential Taxon Abundance Estimations across Ecological Niches

Differential abundance analyses (Wald Test, $P_{\text{val}} = 0.01$) were performed on various ecological models following potential environmental niches suggested by ordination patterns (Figures 8 and 9). Only OTU lineages among comparison groups containing more than 100 sequences were used for each test. We tested the influence of sediment depth in the absence of seepage or hydrothermal influence, the impact of the oxygen minimum zone waters on surficial and subsurface sediments, and the effect of hydrothermal disturbance. These analyses have to be qualified by the fact that they are based on patterns of sequence frequencies, which are derived from the archaeal community but do not necessarily represent it in identical proportions.

To test the estimated differential abundance of archaeal community members in near-surface (depth < 1 mbsf) and subsurface (depth > 1 mbsf) communities under conditions of normal hemipelagic sedimentation, we selected cores ContP03, ContP10, and ContP13; these cores lack hydrothermal, seepage, or OMZ influence and, therefore, show the archaeal community of organic-rich Guaymas Basin sediments in the absence of these selective factors. Here, over 43% of OTUs ($n > 100$ seqs) are differentially abundant with depth (Figures 8A and 8B). Most differentially abundant taxa are estimated to have lower relative abundances in near-surface relative to subsurface control sediment (Figure 8B). Lineages with higher subsurface relative abundances include members of various Bathyarchaeal groups (MCG-1, MCG-3, and MCG-6), a TMEG lineage (OTU 08) closely related to clones previously recovered from Guaymas Basin, and a Hadesarchaea lineage (OTU 18) whose closest relatives are clones from deep Mediterranean waters (Figures 7B and 7C). Among the top 25 OTUs in this study, the only highly abundant lineage that significantly ($P_{\text{val}} = 0.01$) increases in relative abundance in near-surface relative to subsurface control sediment (Figure 8A) is OTU 07, related to Guaymas Basin and Indian estuary sediment MCG-6 clones (Figure 7B). The same lineage was also found to occur preferentially in surficial estuarine sediments in the White Oak River, while avoiding the sulfide-rich, sulfate-reducing, and methanogenic conditions just a few centimeters downcore (Lazar et al., 2015).

When archaeal abundance in all oxygen minimum core OMZP12 samples was checked against the shallow sediment samples of the control cores (ContP03, ContP10, ContP13, see Figures 8C and 8D), over 50% of archaeal OTUs ($n > 100$ seqs) present in surficial controls showed higher relative abundances in core OMZP12. Of the 25 most highly abundant OTUs in this study, those that were differentially abundant in this comparison (14 OTUs) increased in relative abundance in core OMZP12 (eleven of fourteen high

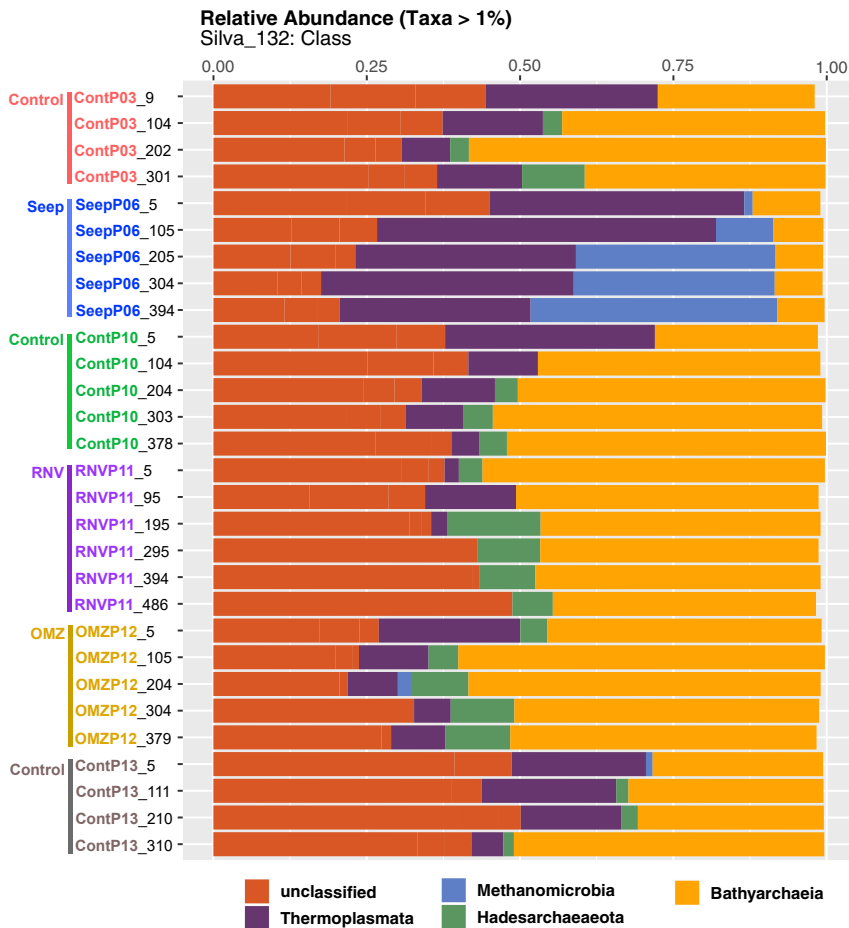


Figure 6. Class-Level Community Composition of All Depths for All Cores in This Study

Core labels are color coded to match the collection sites depicted in the bathymetric map in Figure 1.

abundance OTUs, Figure 8D). Only OTUs 07, 09, and 21, representing the Bathyarchaeal groups MGC-6 and MCG-2, and an MBG-D lineage, respectively, were more abundant in surficial control sediment (Figure 8D).

We also tested for differentially abundant taxa between all core OMZP12 samples and subsurface control sediment (Figure 8E). Here, only 31.1% of OTUs ($n > 100$ seqs) were differentially abundant; of the 25 most abundant OTUs in this study only 4 showed significant abundance differences. Of these four archaeal OTUs, OTUs 15 and 20, Bathyarchaeal lineages in the MCG-1 and MCG-2, respectively, increase in relative abundance in subsurface control sediment. OTU 06 within the MCG-2 group, and OTU 19, an MCG lineage tenuously related (bootstrap value $< 70\%$) to MCG-6, decreased in relative abundance in core OMZP12. Overall, archaeal types occurring in anoxic subsurface sediments of core OMZP12 resemble those in other subsurface cores to a large extent.

The hydrothermally influenced Ringvent core RNVP11 was compared against control core ContP10, located only 1.6 km further west, at the same depth and local sedimentation regime (Figures 9A–9C). In surficial sediment (> 1 mbsf) this comparison revealed only one differentially abundant phylotype, OTU 18, within the Hadesarchaea (Figure 9B). On the other hand, the comparison of subsurface (> 1 mbsf) communities identified 43 OTUs, or 61% of all shared OTUs ($n > 100$ seqs), as differentially abundant between these two sites (Figure 9C). Focusing on the top 25 highest abundance OTUs in this study, eleven OTUs were differentially abundant. Eight taxa, comprising Bathyarchaeal, TMEG, MBG-D, and ANME-1 representatives, were significantly more abundant in subsurface control sediment relative to Ringvent subsurface sediment (OTUs: 07, 08, 10, 11, 14, 15, 16, and 23). The remaining three taxa (OTUs 06, 12, and 25),

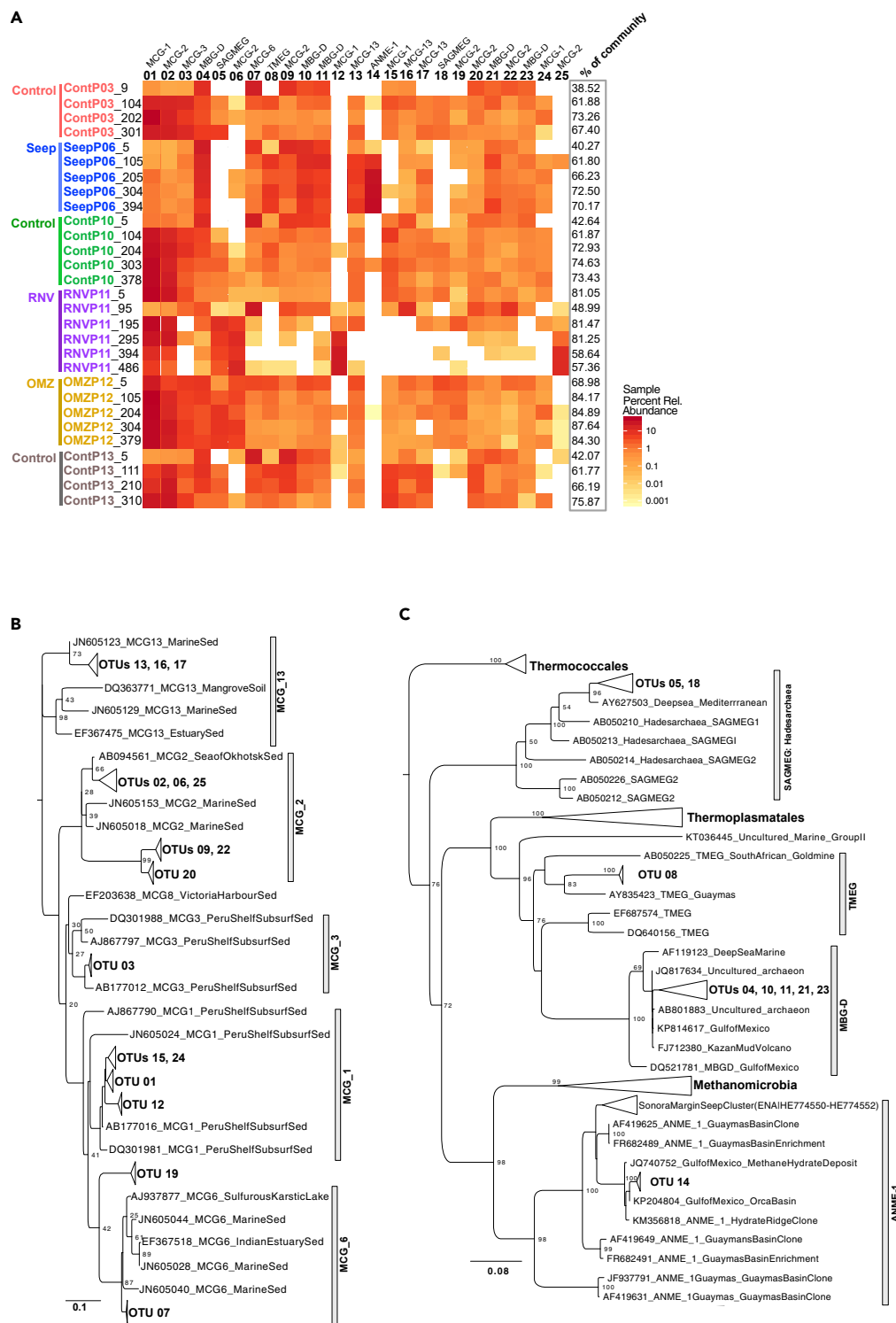


Figure 7. Distribution and Phylogeny of Dominant Archaeal OTUs

(A) Heatmap depicting percent relative abundance distribution for the 25 most abundant OTUs, representing 73.0% of all high-quality sequences in this study, for all cores and across all depths. Core labels are color coded to match the collection sites depicted in the bathymetric map in Figure 1A. The phylogenetic association of each OTU lineage is depicted above each OTU header. The percent of total reads represented by the 25 most abundant OTUs in each

Figure 7. Continued

community is shown in the column labeled “% of community.” Maximum likelihood phylogenetic trees, with 100 bootstrap support, placing the top 25 most abundant OTUs within the following lineages: (B) Bathyarchaea, (C) the Euryarchaeotal lineages MBG-D, TMEG, SAGMEG, and ANMEs.

more abundant in the Ringvent subsurface, were classified as Bathyarchaea related to the MCG-1 and MCG-2 subgroups.

Core SeepP06, from compacted seep sediments, was checked against both shallow and subsurface sediment samples of the control cores (ContP03, ContP10, ContP13, Figures 9D–9F). About 41% of archaeal OTUs ($n > 100$ seqs) present in shallow controls and seep sediment were significantly differentially abundant (Figure 9E). Of the 25 most highly abundant OTUs, most showed higher relative abundances in seep sediment and included lineages classified as ANME-1, MBG-D, and TMEG (Figure 9E). When comparing differentially abundant taxa between core SeepP06 and subsurface control sediment, 53% of archaeal OTUs ($n > 100$ seqs) were differentially abundant (Figure 9F). Most of the OTUs in this comparison, including the most abundant OTUs in this study (OTUs 01–03), increase in relative abundance in control subsurface sediments rather than in core SeepP06 sediment (Figure 9F).

DISCUSSION

Complex Determinants of Archaeal Ecosystem Structure

Overall, complex physical and geochemical factors structure sedimentary habitats and depth-related niches for archaea in Guaymas Basin. Archaeal community ordination patterns reveal niche differentiation and some unexpected clustering patterns among the different sites (Figure 4A). Most notably, surficial communities of background control sediments ContP03, ContP10, and ContP13, at 5–10 cm depth, cluster away from their respective subsurface communities near 1 m depth and below (Figure 4A), consistent with infaunal bioturbation and aeration as observed during Guaymas Basin *Alvin* dives. Taxa co-occurrence network patterns support this differentiation between shallow and seafloor control sediment sites (Figure 5B). The availability of electron acceptors such as oxygen, nitrate, or oxidized metals very likely drives the depth-dependent niche separation observed in background control sediment sites (ContP03, ContP10, and ContP13). Surface archaeal communities continually change as sediment layers accumulate; given high sedimentation rates of 0.23–1 mm/year at Guaymas Basin (Teske et al., 2019), it takes approximately 1,000–4,000 years for background surface communities to transition to subsurface communities at 1 m depth. Interestingly, shallow versus subsurface differentiation is less apparent in seep, OMZ, or hydrothermally influenced sites (Figure 5B); parameters other than sediment depth or surficial redox regime are shaping archaeal community composition in seepage- or hydrothermally influenced habitats, compared with the control sites.

The cores SeepP06 and RNVP11 represent different geochemical regimes (compaction-induced continental margin seepage versus hydrothermal circulation, respectively), yet these two sites cluster tightly in ordination space (Figure 4). Individual geochemical factors, for example, the sulfidic, methane-, and DIC-rich conditions in SeepP06 would have indicated that OMZP12 should be its closest equivalent (Figure 2). The unexpected clustering of SeepP06 and RNVP11 suggests that factors beyond current geochemical conditions, for example, recent environmental disturbance, can influence archaeal community structure. For example, variable ^{14}C age data for the sediment column in core SeepP06 suggested prior perturbation by slumping, for example, during steep slope collapse (Teske et al., 2019). At Ringvent (RNVP11), sedimentary community diversity may have been reduced during prior episodes of thermal purging or high methane flux (Teske et al., 2019), selecting for a resilient, yet potentially less diverse (Figure 3), “survivor” community.

Lastly, community ordination differentiates OMZ sediment from all other sedimentary habitats (Figure 4). Although water depth appears to have a strong influence on OMZ sediment ordination (Figure 4F), we propose that differences in redox potential at the sediment interface due to its direct contact with oxygen-depleted water (Calvert, 1964) rather than water column depth is the key environmental constraint driving the ordination patterns of OMZ sediment. A distinct archaeal community is consistent with the persistence of fully developed oxygen minimum conditions on the Sonora Margin since approximately 13,000 years (Moffitt et al., 2015).

A “Forest View” of Archaea in Guaymas Basin Sediments

Archaea observed in this sedimentary habitat survey belong to the Bathyarchaea, the MBG-D and TMEG lineages within the Thermoplasmatales, the Hadesarchaea (SAGMEG), and ANME-1 lineages, as shown

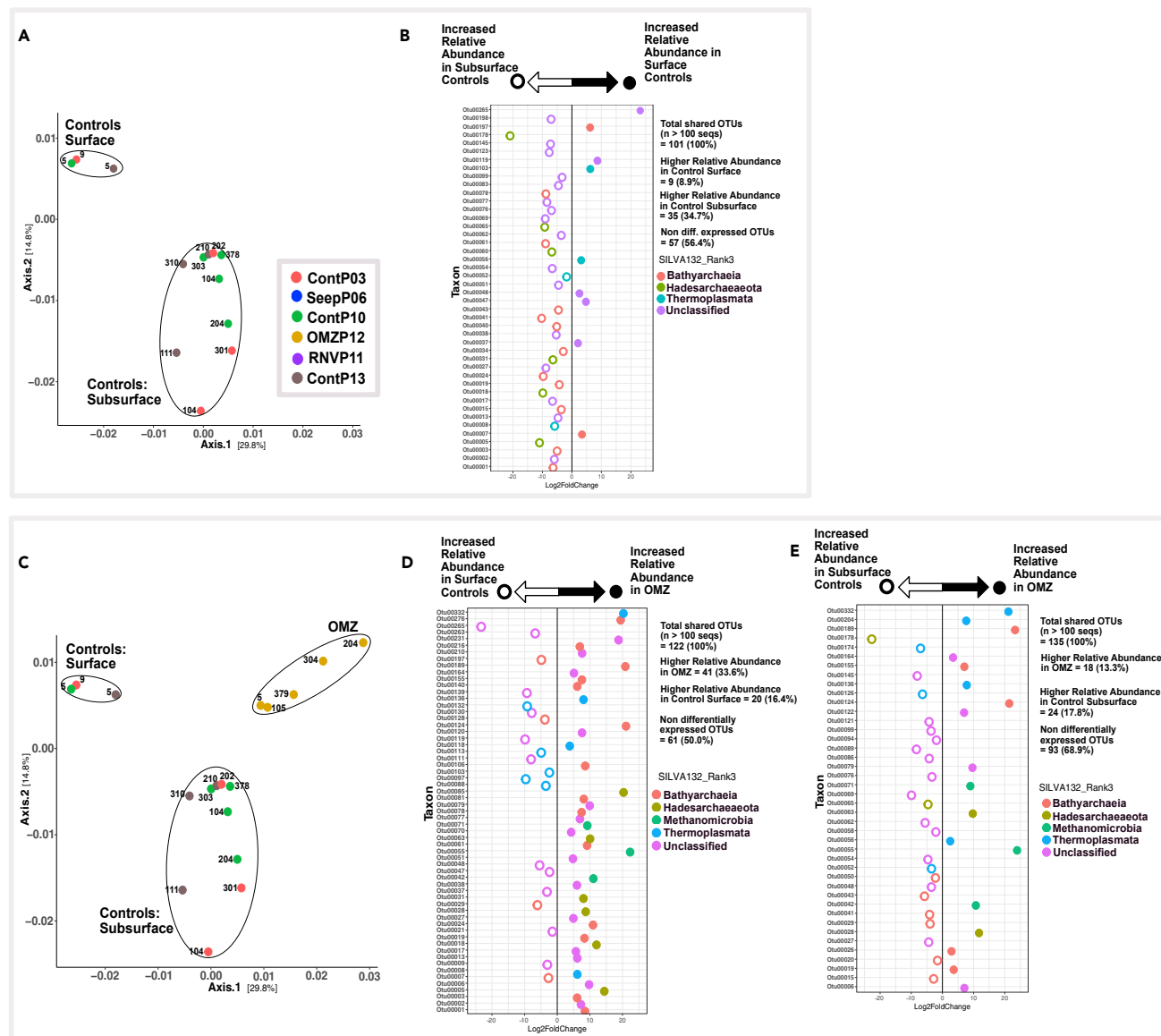


Figure 8. Differential Abundance Analyses Based on Wald's Test (Significance: alpha = 0.01)

(A) Ordination depicting archaeal community clustering for surface and subsurface samples of control sites ContP03, ContP10, and ContP13.

(B) Differentially abundant OTUs in near-surface versus subsurface communities from control sites.

(C) Ordination depicting community clustering in OMZP12, and surficial versus subseafloor control sites.

(D and E) Differentially abundant OTUs in OMZP12 compared with surficial and (E) subsurface communities from control sites. Note: OTUs 13, 17, and 21 are color coded as "unclassified" by SILVA132, but phylogeny analysis (see Figures 7B and 7D) places them as members of MCG-13 (OTUs 13 and 17) and MBG-D (OTU 21).

previously in a sequencing survey using different archaeal primers (Teske et al., 2019). The uncultured Bathyarchaea and MBG-D archaea pronouncedly dominate the dataset, and cold anoxic marine sediments globally (Kubo et al., 2012; Lloyd et al., 2013). Bathyarchaea play an important role, tantamount to that of the domain Bacteria, in the remineralization of complex organic matter in marine sediment (Lloyd et al., 2013; Orsi et al., 2020); some of their members (MCG-8 lineage) use lignin, the second most common biopolymer on Earth, as an energy source (Yu et al., 2018). Since Bathyarchaeota harbor the Wood-Ljungdahl pathway (WLP), they are implied in acetogenic subsurface metabolism (He et al., 2016). Recycling fermentation products (CO₂, hydrogen) with the acetogenic WLP during the breakdown of diverse substrate types may provide the Bathyarchaea with an energetic advantage over classical fermenters in anoxic

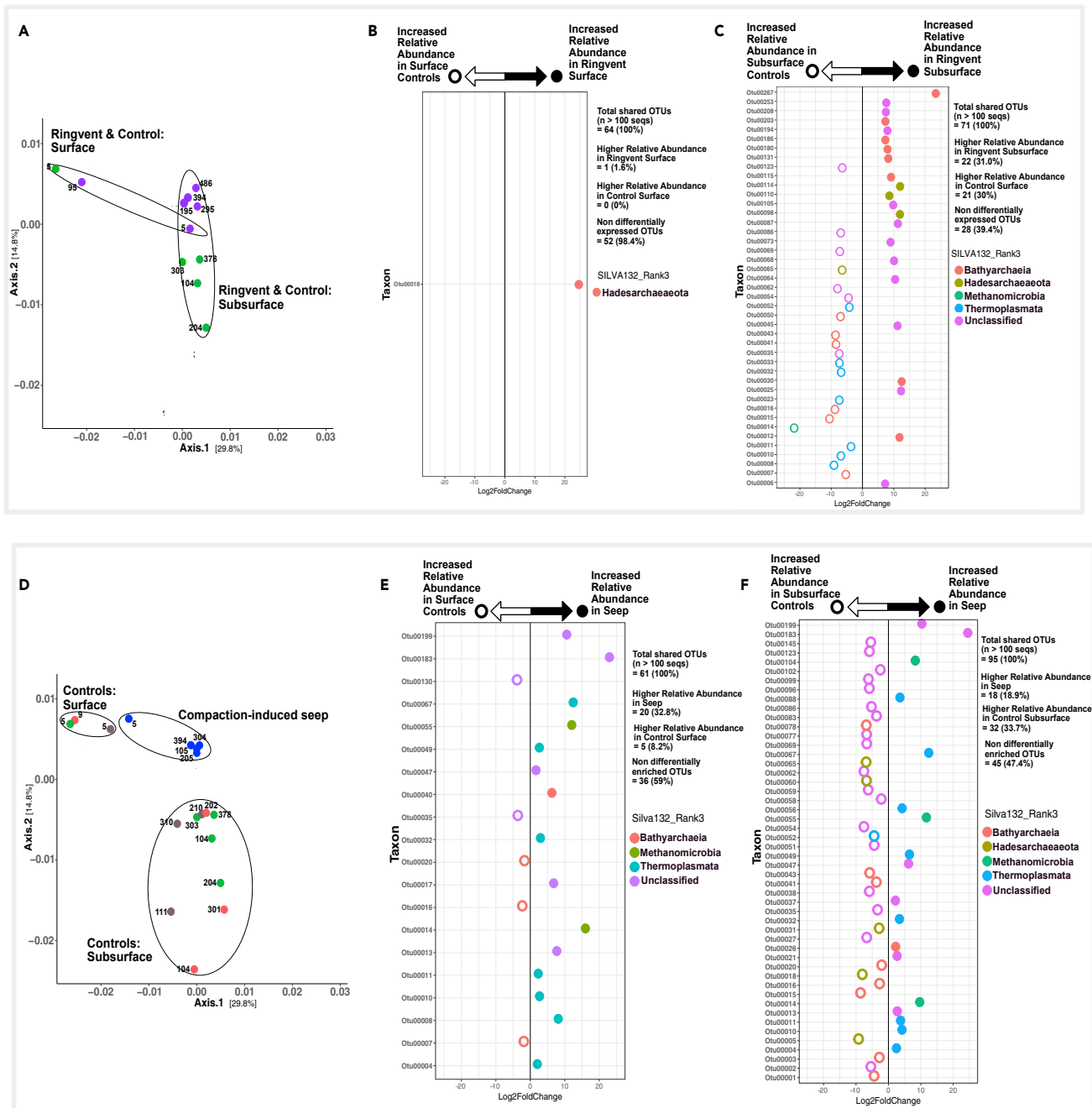


Figure 9. Differential Abundance Analyses Based on Wald's Test (Significance: alpha = 0.01)

(A–C) (A) Ordination depicting community clustering in RNVP11 and ContP10. Differentially abundant OTUs in RNVP11 and ContP10 for (B) surface samples and (C) subsurface samples.

(D) Ordination depicting community clustering in SeepP06, and surficial and seafloor control sites.

(E and F) Differentially abundant OTUs in SeepP06 samples compared with (E) surficial and (F) subsurface communities from control sites. Note: OTUs 13, 17, and 21 are color coded as “unclassified” by SILVA132, but phylogeny analysis (see [Figures 7B and 7D](#)) places them as members of MCG-13 (OTUs 13 and 17) and MBG-D (OTU 21).

sediment ([Schuchmann and Müller, 2016](#)). It remains open whether acetogenic pathways are used for net autotrophy or derive their substrates primarily from organic carbon sources ([Lever et al., 2010](#)); similar considerations apply to the metabolically versatile MBG-D archaea ([Zhou et al., 2019](#)). Some Bathyarchaea harbor genes of the MCR complex, suggesting methylotrophic methanogenic activity and, perhaps,

syntrophic interactions with sulfate-reducing bacteria leading to the anaerobic oxidation of methane (Evans et al., 2015). However, the dominant Bathyarchaea OTUs recovered in this study (MCG groups 1, 2, 3, 6, and 13, Figure 7B) are only distant relatives of methane-cycling marine Bathyarchaea, which fall into MCG groups 15 and 16 (Evans et al., 2015). Hadesarchaea, originally described as the South-African Gold Mine Miscellaneous Euryarchaeal Group (SAGMEG), are metabolically versatile anaerobic heterotrophs with the metabolic potential for CO and H₂ oxidation coupled with nitrite reduction to ammonia and are found in environments across broad (4°C –80°C) thermal ranges (Baker et al., 2016). One of two frequently recovered Hadesarchaea lineages (OTU 05) is conspicuously present in high relative abundance in subsurface sediment in Ringvent (RNVP11), where low observed sequence richness (Figure 3) coincides with evidence (high silica porewater concentrations at depth indicative of a thermal dissolution of sedimentary diatoms, Figure 2I) for a thermal purge in the past (Teske et al., 2019).

Unsurprisingly, the methane-cycling Methanomicrobia are rare in background sediments (ContP03, Cont10, and Cont13) but increase in relative abundance in core SeepP06 and, to a much lower extent (slightly over 2%), in core OMZP12 at 204 cm depth (Figures 6 and S2). Interestingly, a single ANME-1 OTU lineage, OTU14, is highly abundant in SeepP06 sediments (Figures 7A and 7C). This lineage is closely related to ANMEs recovered from cold, anoxic habitats, such as seafloor seep sediments, methane hydrates, and hypersaline anoxic basins, and distinct from previously described ANME-1 phylotypes from Sonora Margin cold seep sediments and potentially thermophilic ANME-1 phylotypes from Guaymas Basin hydrothermal sediments (Biddle et al., 2012; Holler et al., 2011). Although ANME-1 archaea were generally assumed to be obligate methanotrophs, this assumption has been challenged and this lineage has been proposed as potentially methanogenic, based on its occurrence and activity in sulfate-depleted sediments (Lloyd et al., 2011; Kevorkian et al., 2020); thus, the biogeochemical role of these archaea would be modulated by the presence or absence of sulfate or concomitant changes in electron donors. ANME-2 and cultured methanogenic lineages were observed in low percent abundances in all cores in this study (Figure S2). Interestingly, ANME-2 lineages were extremely rare, representing less than 0.05% of any sample and less than 0.02% of any SeepP06 community (Figure S2, Table S3). The prevalence of ANME-1 over ANME-2 in our survey is consistent with the ecophysiological preference of ANME-1 archaea for reducing, sulfidic subsurface sediments and the preference of ANME-2 for near-surface sediments with intermittently oxidizing conditions (Ruff et al., 2015; Rossel et al., 2011). Previous surveys of mat-covered seep sediments on the Sonora Margin have revealed transitions from ANME-2 toward ANME-1 within short push cores of maximum 17 cm depth (Vigneron et al., 2013).

Overall, we hypothesize that the archaeome in the sedimented flanking regions of Guaymas Basin is generally fueled by heterotrophic processes including the degradation of proteins, polymeric carbohydrates (Ziervogel and Arnosti, 2020), and accumulating lipids (Teske et al., 2002) resulting from high sedimentation rates driven by high levels of primary production in the water column. Diverse niche communities allow the Guaymas archaeome to adapt to environmental challenges, such as hydrothermalism or methane seepage, that are common in the greater Guaymas Basin area.

Ecological Comparisons: Differentially Abundant Taxa across Sedimentary Habitats

Near-Surface versus Subsurface Sediment Niches

When comparing the surficial to the subsurface archaeal populations in background control sediments, the majority of OTUs estimated to be differentially abundant are significantly more abundant in the subsurface relative to the surficial sediment, suggesting that benthic archaea prefer subsurface conditions (Figures 8A and 8B). This trend may also reflect the impact of electron acceptors; for example, oxygen permeates background sediments in Guaymas Basin for at least 1 cm (Teske et al., 2016, Figure 8B therein). Following a recently proposed model for benthic microbial communities (Starnawski et al., 2017) the archaeal community at the oxic water-sediment interface likely undergoes downcore selection, based on site-specific selective pressure, resulting in reduced diversity with depth but a higher prevalence of subsurface-adapted taxa within a few thousand years after burial in anoxic subseafloor sediment. Benthic archaea, predominantly Bathyarchaea and MBG-D lineages, survive on residual carbon sources that remain after burial and microbial degradation in surficial sediments (Lloyd et al., 2013). Interestingly, catabolic activity and electron donor diversity, rather than terminal electron acceptor type or burial time, appear to drive bacterial OTU richness in anoxic subseafloor sediment (Walsh et al., 2016). This niche construction mechanism, driven by the biotic microenvironment as opposed to abiotic environmental filtering (Aguilar-Trigueros

et al., 2017), is potentially widespread across the large habitable volume represented by non-hydrothermally influenced seafloor sediments in Guaymas Basin.

OMZ versus Control Sediment

When comparing OMZ and surficial background control sedimentary communities, 50% of high abundance OTUs found across both habitats are significantly differentially abundant (Figure 8D). Two-thirds of the differentially abundant taxa have higher relative abundances in the OMZ rather than the surficial background sediments. The MCG lineages MGC-6 and MCG-2 (OTUs 9 and 7) and an MBG-D phylotype (OTU 21) increase in relative abundance in the surficial background controls relative to the OMZ sediment (Figure 8D). Interestingly, MGC-6 members bear hydrolases that specifically target plant-derived polymeric carbohydrates (Lazar et al., 2016), a potential trait-environment relationship that may differentiate surficial background control from OMZ sediment communities (Figure 8D). When comparing subsurface background control and OMZ sediment communities, the number of differentially abundant OTUs was about equal across both environments; however, the majority (68.9%) of high abundance OTUs show no significant differences in differential abundances across subsurface background controls and OMZ environments (Figure 8E). This implies that the subsurface, rather than surficial, background control communities are more similar to the OMZ communities, a point also corroborated by taxa co-occurrence network analysis (Figure 5B). Thus, oxygen depletion in background subsurface sediment and oxygen depletion through the overlying oxygen minimum zone of the water column (Calvert, 1964) result in some convergence between archaeal communities across geographically distant and environmentally distinct sedimentary habitats.

Ringvent versus Control Sediment

The surficial archaeal communities of Ringvent (RNVP11) and its nearby control site (ContP10) are similar to each other, as indicated by extensive co-occurrence networks (Figure 5) and by the dearth of differentially abundant OTUs between the two cores (Figure 9B). A member of the Hadesarchaea, OTU18, is the only differentially abundant lineage in Ringvent surficial sediment relative to the control; otherwise differences in taxon relative abundance across these habitats are negligible. These sites are only 1.6 km apart and therefore most likely share recent depositional histories and microbial inoculum sources, which validates core ContP10 as a site-specific control for assessing the environmental determinants structuring subsurface archaeal communities at Ringvent. The reduction in sequence recovery and, potentially, archaeal community richness in subsurface Ringvent (RNVP11) sediment (Figure 3) is attributed to environmental selection via hydrothermal purging as reflected in silica dissolution, or methane seepage driven by recent sill emplacement that continues to drive hydrothermal circulation, selecting against microbes unable to withstand these chemical or thermal changes (Teske et al., 2019). Thus, OTUs with increased relative abundances in Ringvent subsurface sediment compared with its nearby control site (Figure 9C) may occur via two possible ecological scenarios: (1) surviving resilient microbes could dominate the habitat after their competitors have been removed and (2) new arrivals after the disturbance could efficiently recolonize the depopulated surface sediment.

Seep versus Control Sediment

Differential abundance comparisons show that the ANME-1, MBG-D, and TMEG lineages increase in relative abundance in the seep sediments, compared with controls (Figures 9D and 9E). Generally, methane seeps are specialized microbial benthic habitats where methanotrophic archaea (ANME) and syntrophic Deltaproteobacteria oxidize methane anaerobically exploiting sulfate as an electron acceptor (Lloyd et al., 2010; Ruff et al., 2015). The dominance of these inter-domain syntrophic partners distinguishes seafloor seep habitats (Ruff et al., 2015). Archaeal community structure in SeepP06 sediments differs little with depth; it is most similar, in terms of taxa overlap, to other samples from the same core (Figures 4 and 5). Therefore, the influence of cold seepage drives community selection to a greater degree than the environmental factors associated with depth-dependent niche differentiation observed in background control sediment.

Comparison with other Sonora Margin cores highlights the seep characteristics of core SeepP06. Based on the presence or absence of major archaeal lineages, SeepP06 archaeal communities are similar to surficial (<1 mbsf) communities from Sonora Margin cold seeps, predominantly comprising Thermoplasmata (MBG-D), Bathyarchaea, and ANME lineages (Cruaud et al., 2017). The SeepP06 archaeal communities share dominant archaeal lineages—the Thermoplasmatales (MBG-D), Lokiarchaeota, and

Bathyarchaeota—with Sonora Margin subsurface sediments (core BCK1 [Vigneron et al., 2014]). Interestingly, the high proportion of ANME-1 archaea in SeepP06 is not shared by the Sonora Margin subsurface core (Vigneron et al., 2014). The Sonora Margin subsurface sediment core has a deeper methane/sulfate interface than SeepP06, ca. 4–5 m instead of 1 m, and contains little sulfide above 5 m depth, indicating strongly attenuated seep influence in core BCK1 compared with SeepP06.

Core-Specific Features of the Benthic Archaeome

Controls that structure microbial communities in hydrothermal sediments of Guaymas Basin have been studied extensively; for example, extreme temperature and porewater gradients shape microbial population structure, genomic repertoire, and activities within a few centimeters depth beneath the seafloor (McKay et al., 2012, 2016; Dombrowski et al., 2018). However, ecological factors influencing microbial life in other sedimentary habitats at Guaymas Basin are comparatively unconstrained. By comparing archaeal communities in diverse sedimentary habitats to background controls representative of standard hemipelagic sedimentation, characteristic responses of the archaeal communities to these distinct environmental settings are becoming apparent. Compaction-induced seepage near the base of the Sonora Margin, and the resulting methane- and sulfide-rich porewater conditions in core SeepP06, selected for anaerobic methane-oxidizing archaea (ANME-1) and for MBG-D archaea within the Thermoplasmata, and reduced the relative proportion of Hadesarchaea and Bathyarchaeota. Prior disturbances by hydrothermal impact or strong methane seepage, exemplified in the Ringvent sediments (RNVP11), also strongly differentiated sedimentary archaeal communities from those in background controls. Observed community richness in RNVP11 based on rarefaction curves appears reduced throughout much of the core; these results resembled the outcome of a parallel study using different archaeal primers, and bacterial primers as well (Teske et al., 2019). Lastly, anoxic bottom waters impinging on the sediment on the upper Sonora Margin (OMZP12) drive similarities between anoxic surficial sediment at this site and anoxic subsurface background control sediments. The anoxic redox state of the water-sediment interface may also enhance archaeal richness estimates in the upper sediment column, potentially by facilitating the pelagic-benthic transition of archaea or selecting against a bacteria-dominated interface (Xia et al., 2017). In brief, the archaeal communities of different cores respond in different ways to specific local controls.

Environmental History Determines Ecological Context

The sediment cores shared similar biogeochemical parameters, such as sedimentary TOC, and organic matter $\delta^{13}\text{C}$, $\delta^{15}\text{N}$ and C:N ratios. Repeatedly, studies of uncultured microbes in the sedimentary subsurface tried to correlate community composition with a wide range of biogeochemical or thermal parameters, in the hope that these linkages provide insights into habitat preference and ecophysiology of uncultured archaea (Lazar et al., 2015; McKay et al., 2016; Durbin and Teske, 2012). Although this strategy can yield valuable results, we caution that patterns of archaeal community composition are not deterministically linked to biogeochemical parameters alone, rather, the full context of an ecological interpretation requires that biological and geochemical observations are integrated with the environmental setting and history of a site. For example, the lighter $\delta^{13}\text{C}$ values of sedimentary organic matter in RNVP11 (trending toward -22‰ compared with most values clustering between 20‰ and 21‰), the slightly elevated C:N ratios at this site, increased Si concentrations at depth, or the elevated methane content superimposed on seawater-like porewater characteristics are not in themselves critical factors that determine biological metrics in this core; these factors are significant because they reveal a depositional history of organic-rich sediments overprinted by relatively recent hydrothermalism and methane flux that has left its footprint on the present-day archaeal community. In another example, the archaeal communities of cores SeepP06 and OMZP12 would be assumed to be similar, since both sites are rich in sulfide, methane, and DIC and show rapid sulfate depletion concomitant with methane accumulation; these characteristics indicate strongly reducing marine seep-like conditions with microbial methane and sulfur cycling by functionally equivalent microbial communities. However, the distinct environmental settings and histories of these two cores, at the heavily compacted, seep-influenced base of the Sonora Margin (SeepP06), and under the persistent oxygen minimum zone waters of the upper Sonora Margin (OMZP12), ultimately select for different archaeal communities.

Conclusion

In the greater Guaymas Basin and Sonora Margin area, complex geological and oceanographic processes impose environmental controls on different sedimentary habitats and their archaeal populations relative to background control sites. In background sediments, archaeal communities vary little with depth after the

surface/subsurface transition; here, subsurface communities result primarily from long-term survival likely conferred by relatively reduced mortality (Kirkpatrick et al., 2019). In contrast, localized factors, including water column anoxia, methane seepage, and hydrothermal circulation, constrain the biodiversity and potential biogeochemical activity of sedimentary Archaea across our benthic survey in specific ways. Overall, our observations suggest that local sediment biogeochemistry should be viewed in a broader context—within the history and evolution of a particular site—to reveal its influence on selective survival for certain lineages and subsequent shaping of the resident archaeal community.

Limitations of the Study

The compositional nature of amplicon-based sequence studies precludes discussion of absolute abundances. Piston coring disturbs small-scale structures near the water-sediment interface; thus, the potential effect of bioturbation on surficial sediment redox state is not addressed in this study.

Resource Availability

Lead Contact

Gustavo A. Ramírez: ramirezg@westernu.edu.

Materials Availability

This study did not generate new unique reagents.

Data and Code Availability

All sequence data are publicly available at the following repository: NCBI under BioProject PRJNA553578 and accession numbers SRX6444849–SRX6444877.

Geochemical data are available at the BCO-DCO under these reference links:

Porewater methane data: <https://www.bco-dmo.org/dataset/661750/data>.

Porewater sulfate data: <https://www.bco-dmo.org/dataset/661775/data>.

Porewater DIC data: <https://www.bco-dmo.org/dataset/661658/data>.

Porewater sulfide: <https://www.bco-dmo.org/dataset/661808/data>.

METHODS

All methods can be found in the accompanying [Transparent Methods supplemental file](#).

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.isci.2020.101459>.

ACKNOWLEDGMENTS

Sampling in Guaymas Basin was funded by NSF OCE grant 1449604 “Rapid Proposal: Guaymas Basin site survey cruise for IODP proposal 833” to A.P.T.; NSF C-DEBI grant “Characterizing seafloor life and environments in Guaymas Basin” to A.P.T. and A.C.R. This is C-DEBI Publication No. 542. Thanks go to Zachary Stewart for helping with the sedimentary organic matter geochemistry analyses. We thank the Science crew of R/V *El Puma* for excellent piston coring skills and a very enjoyable cruise.

AUTHOR CONTRIBUTIONS

A.P.T. conceived the study. L.J.M., A.C.R., and A.P.T. collected samples. C.M. led the *El Puma* cruise and the piston coring effort. A.B. extracted DNA from sediments. L.J.M. performed sequencing in the laboratory of M.W.F. A.C.R., L.J.M., and C.H. performed biogeochemical and sedimentological analyses. G.A.R. performed bioinformatic analyses with phylogeny input from A.P.T. G.A.R. and A.P.T. wrote the manuscript with input from all authors.

DECLARATION OF INTERESTS

There are no competing interests.

Received: June 5, 2020

Revised: July 31, 2020

Accepted: August 11, 2020

Published: September 25, 2020

REFERENCES

- Aguilar-Trigueros, C.A., Rillig, M.C., and Ballhausen, M.-B. (2017). Environmental filtering is a relic. A response to cadotte and tucker. *Trends Ecol. Evol.* **32**, 882–884.
- Baker, B.J., Saw, J.H., Lind, A.E., Lazar, C.S., Hinrichs, K.U., Teske, A.P., and Ettema, T.J. (2016). Genomic inference of the metabolism of cosmopolitan subsurface Archaea, Hadesarchaea. *Nat. Microbiol.* **1**, 16002.
- Biddle, J.F., Cardman, Z., Mendlovitz, H., Albert, D.B., Lloyd, K.G., Boetius, A., and Teske, A. (2012). Anaerobic oxidation of methane at different temperature regimes in Guaymas Basin hydrothermal sediments. *ISME J.* **6**, 1018–1031.
- Calvert, S.E. (1964). Factors affecting distribution of laminated diatomaceous sediments in the Gulf of California. In *Marine Geology of the Gulf of California, Vol. 3*, T.H. van Andel and G.G. Shor, eds. (American Association of Petroleum Geologists Memoir), pp. 311–330.
- Calvert, S.E. (1966). Origin of diatom-rich, varved sediments from the Gulf of California. *J. Geol.* **74**, 546–565.
- Campbell, A.C., and Gieskes, J.M. (1984). Water column anomalies associated with hydrothermal activity in the Guaymas Basin, Gulf of California. *Earth Planet. Sci. Lett.* **68**, 57–72.
- Cruaud, P., Vigneron, A., Pignet, P., Caprais, J.-C., Lesongeur, F., Toffin, L., Godfroy, A., and Cambon-Bonavita, M.-A. (2017). Comparative study of Guaymas basin microbiomes: cold seeps vs. hydrothermal vents sediments. *Front. Mar. Sci.* **4**, 417.
- Didy, B.M., and Simoneit, B.R.T. (1989). Hydrothermal oil of Guaymas Basin and implications for petroleum formation mechanisms. *Nature* **342**, 65–69.
- Dombrowski, N., Teske, A., and Baker, B.J. (2018). Exapnsive microbial metabolic versatility and biodiversity in dynamic Guaymas Basin hydrothermal sediments. *Nat. Commun.* **9**, 4999.
- Durbin, A.M., and Teske, A. (2012). Archaea in organic-lean and organic-rich marine subsurface sediments: an environmental gradient reflected in distinct phylogenetic lineages. *Front. Microbiol.* **3**, 168.
- Evans, P., Parks, D.H., Chadwick, G.L., Robbins, S.J., Orphan, V.J., Golding, S.D., and Tyson, G.W. (2015). Methane metabolism in the archaeal phylum Bathyarchaeota revealed by genome-centric metagenomics. *Science* **350**, 434–438.
- Geilert, S., Hensen, C., Schmidt, M., Liebetrau, V., Scholz, F., Doll, M., Deng, L., Fiskal, A., Lever, M.A., Su, C.-C., et al. (2018). On the formation of hydrothermal vents and cold seeps in the Guaymas Basin, Gulf of California. *Biogeosciences* **15**, 5715–5731.
- He, Y., Li, M., Perumal, V., Feng, X., Fang, J., Xie, J., Sievert, S.M., and Wang, F. (2016). Genomic and enzymatic evidence for acetogenesis among multiple lineages of the archaeal phylum Bathyarchaeota widespread in marine sediments. *Nat. Microbiol.* **1**, 16035.
- Holler, T., Widdel, F., Knittel, K., Amann, R., Kellermann, M.Y., Hinrichs, K.U., Teske, A., Boetius, A., and Wegener, G. (2011). Thermophilic anaerobic oxidation of methane by marine microbial consortia. *ISME J.* **5**, 1946–1956.
- Kevorkian, R., Callahan, S., Winstead, R., and Lloyd, K.G. (2020). ANME-1 archaea drive methane accumulation and removal in estuarine sediment. *bioRxiv*. <https://doi.org/10.1101/2020.02.24.963215>.
- Kirkpatrick, J.B., Walsh, E.A., and D'hondt, S. (2019). Microbial selection and survival in subseafloor sediment. *Front. Microbiol.* **10**, 956.
- Kubo, K., Lloyd, K.G., Biddle, J.F., Amann, R., Teske, A., and Knittel, K. (2012). Archaea of the Miscellaneous Crenarchaeotal Group are abundant, diverse and widespread in marine sediments. *ISME J.* **6**, 1949–1965.
- Lazar, C.S., Baker, B.J., Seitz, K., Hyde, A.S., Dick, G.J., Hinrichs, K.U., and Teske, A.P. (2016). Genomic evidence for distinct carbon substrate preferences and ecological niches of Bathyarchaeota in estuarine sediments. *Environ. Microbiol.* **18**, 1200–1211.
- Lazar, C.S., Biddle, J.F., Meador, T.B., Blair, N., Hinrichs, K.U., and Teske, A.P. (2015). Environmental controls on intragroup diversity of the uncultured benthic archaea of the miscellaneous Crenarchaeotal group lineage naturally enriched in anoxic sediments of the White Oak River estuary (North Carolina, USA). *Environ. Microbiol.* **17**, 2228–2238.
- Lever, M.A., Heuer, V.B., Morono, Y., Masui, N., Schmidt, F., Alperin, M.J., Inagaki, F., Hinrichs, K.-U., and Teske, A. (2010). Acetogenesis in deep subseafloor sediments of the Juan de Fuca Ridge Flank: a synthesis of geochemical, thermodynamic, and gene-based evidence. *Geomicrobiol. J.* **27**, 183–211.
- Lizarralde, D., Axen, G.J., Brown, H.E., Fletcher, J.M., Gonzalez-Fernandez, A., Harding, A.J., Holbrook, W.S., Kent, G.M., Paramo, P., Sutherland, F., and Umhoefer, P.J. (2007). Variation in styles of rifting in the Gulf of California. *Nature* **448**, 466–469.
- Lizarralde, D., Soule, S.A., Seewald, J.S., and Proskurowski, G. (2010). Carbon release by off-axis magmatism in a young sedimented spreading centre. *Nat. Geosci.* **4**, 50–54.
- Lloyd, K.G., Albert, D.B., Biddle, J.F., Chanton, J.P., Pizarro, O., and Teske, A. (2010). Spatial structure and activity of sedimentary microbial communities underlying a *Beggiatoa* spp. mat in a Gulf of Mexico hydrocarbon seep. *PLoS One* **5**, e8738.
- Lloyd, K.G., Alperin, M.J., and Teske, A. (2011). Environmental evidence for net methane production and oxidation in putative ANAerobic METHanotrophic (ANME) archaea. *Environ. Microbiol.* **13**, 2548–2564.
- Lloyd, K.G., Schreiber, L., Petersen, D.G., Kjeldsen, K.U., Lever, M.A., Steen, A.D., Stepanauskas, R., Richter, M., Kleindienst, S., Lenk, S., et al. (2013). Predominant archaea in marine sediments degrade detrital proteins. *Nature* **496**, 215–218.
- Lonsdale, P., and Becker, K. (1985). Hydrothermal plumes, hot springs, and conductive heat flow in the Southern Trough of Guaymas Basin. *Earth Planet. Sci. Lett.* **73**, 211–225.
- Martens, C.S. (1990). Generation of short chain organic acid anions in hydrothermally altered sediments of the Guaymas Basin, Gulf of California. *Appl. Geochem.* **5**, 71–76.
- McKay, L., Klokman, V.W., Mendlovitz, H.P., Larowe, D.E., Hoer, D.R., Albert, D., Amend, J.P., and Teske, A. (2016). Thermal and geochemical influences on microbial biogeography in the hydrothermal sediments of Guaymas Basin, Gulf of California. *Environ. Microbiol. Rep.* **8**, 150–161.
- McKay, L.J., Macgregor, B.J., Biddle, J.F., Albert, D.B., Mendlovitz, H.P., Hoer, D.R., Lipp, J.S., Lloyd, K.G., and Teske, A.P. (2012). Spatial heterogeneity and underlying geochemistry of phylogenetically diverse orange and white *Beggiatoa* mats in Guaymas Basin hydrothermal sediments. *Deep Sea Res.* **67**, 21–31.
- McMurdie, P.J., and Holmes, S. (2014). Waste not, want not: why rarefying microbiome data is inadmissible. *PLoS Comput. Biol.* **10**, 4.
- Moffitt, S.E., Moffitt, R.A., Sauthoff, W., Davis, C.V., Hewett, K., and Hill, T.M. (2015). Paleooceanographic insights on recent oxygen minimum zone expansion: lessons for modern oceanography. *PLoS One* **10**, e0115246.
- Orsi, W.D., Vuillemin, A., Rodriguez, P., Coskun, O.K., Gomez-Saez, G.V., Lavik, G., Mohrholz, V., and Ferdelman, T.G. (2020). Metabolic activity analyses demonstrate that Lokiarchaeon exhibits

- homoacetogenesis in sulfidic marine sediments. *Nat. Microbiol.* 5, 248–255.
- Paull, C.K., Ussler, W., Peltzer, E.T., Brewer, P.G., Keaten, R., Mitts, P.J., Nealon, J.W., Greinert, J., Herguera, J.-C., and Elena Perez, M. (2007). Authigenic carbon entombed in methane-soaked sediments from the northeastern transform margin of the Guaymas Basin, Gulf of California. *Deep Sea Res.* 54, 1240–1267.
- Peter, J.M., Peltonen, P., Scott, S.D., Simoneit, B.R.T., and Kawka, O.E. (1991). 14C ages of hydrothermal petroleum and carbonate in Guaymas Basin, Gulf of California: implications for oil generation, expulsion, and migration. *Geology* 19, 253–256.
- Peter, J.M., and Scott, S.D. (1988). Mineralogy, composition, and fluid-inclusion microthermometry of seafloor hydrothermal deposits in the southern trough of Guaymas Basin, Gulf of California. *Can. Mineral.* 26, 567–587.
- Portail, M., Olu, K., Escobar-Briones, E., Caprais, J.C., Menot, L., Waeles, M., Cruaud, P., Sarradin, P.M., Godfroy, A., and Sarrazin, J. (2015). Comparative study of vent and seep macrofaunal communities in the Guaymas Basin. *Biogeosciences* 12, 5455–5479.
- Rossel, P.E., Elvert, M., Ramette, A., Boetius, A., and Hinrichs, K.-U. (2011). Factors controlling the distribution of anaerobic methanotrophic communities in marine environments: evidence from intact polar membrane lipids. *Geochim. Cosmochim. Acta* 75, 164–184.
- Ruff, S.E., Biddle, J.F., Teske, A.P., Knittel, K., Boetius, A., and Ramette, A. (2015). Global dispersion and local diversification of the methane seep microbiome. *Proc. Natl. Acad. Sci. U S A* 112, 4015–4020.
- Schuchmann, K., and Müller, V. (2016). Energetics and Application of heterotrophy in acetogenic bacteria. *Appl. Environ. Microbiol.* 82, 4056–4069.
- Seewald, J.S., Seyfried, W.E., and Thornton, E.C. (1990). Organic-rich sediment alteration: an experimental and theoretical study at elevated temperatures and pressures. *Appl. Geochem.* 5, 193–209.
- Simoneit, B.R.T., Lonsdale, P., Edmond, J.M., and Shank, W.C., III (1990). Deep-water hydrocarbon seeps in Guaymas Basin, Gulf of California. *Appl. Geochem.* 5, 41–49.
- Spang, A., Caceres, E.F., and Ettema, T.J.G. (2017). Genomic exploration of the diversity, ecology, and evolution of the archaeal domain of life. *Science* 357, eaaf3883.
- Starnawski, P., Bataillon, T., Ettema, T.J., Jochum, L.M., Schreiber, L., Chen, X., Lever, M.A., Polz, M.F., Jorgensen, B.B., Schramm, A., and Kjeldsen, K.U. (2017). Microbial community assembly and evolution in subseafloor sediment. *Proc. Natl. Acad. Sci. U S A* 114, 2940–2945.
- Teske, A., De Beer, D., McKay, L.J., Tivey, M.K., Biddle, J.F., Hoer, D., Lloyd, K.G., Lever, M.A., Roy, H., Albert, D.B., et al. (2016). The Guaymas basin hiking guide to hydrothermal mounds, chimneys, and microbial mats: complex seafloor expressions of subsurface hydrothermal circulation. *Front. Microbiol.* 7, 75.
- Teske, A., Hinrichs, K.U., Edgcomb, V., De Vera Gomez, A., Kysela, D., Sylva, S.P., Sogin, M.L., and Jannasch, H.W. (2002). Microbial diversity of hydrothermal sediments in the Guaymas Basin: evidence for anaerobic methanotrophic communities. *Appl. Environ. Microbiol.* 68, 1994–2007.
- Teske, A., McKay, L., Ravelo, A.C., Aiello, I., Mortera, C., Núñez-Useche, F., Canet, C., Chanton, J.P., Brunner, B., Hensen, C., et al. (2019). Characteristics and evolution of sill-driven off-axis hydrothermalism in Guaymas Basin—the Ringvent site. *Sci. Rep.* 9, 13847.
- Vigneron, A., Cruaud, P., Pignet, P., Caprais, J.C., Cambon-Bonavita, M.A., Godfroy, A., and Toffin, L. (2013). Archaeal and anaerobic methane oxidizer communities in the Sonora Margin cold seeps, Guaymas Basin (Gulf of California). *ISME J.* 7, 1595–1608.
- Vigneron, A., Cruaud, P., Roussel, E.G., Pignet, P., Caprais, J.C., Callac, N., Ciobanu, M.C., Godfroy, A., Cragg, B.A., Parkes, J.R., et al. (2014). Phylogenetic and functional diversity of microbial communities associated with subsurface sediments of the Sonora Margin, Guaymas Basin. *PLoS One* 9, e104427.
- Von Damm, K.L., Edmond, J.M., Measures, C.I., and Grant, B. (1985). Chemistry of submarine hydrothermal solutions at Guaymas Basin, Gulf of California. *Geochim. Cosmochim. Acta* 49, 2221–2237.
- Walsh, E.A., Kirkpatrick, J.B., Pockalny, R., Sauvage, J., Spivack, A.J., Murray, R.W., Sogin, M.L., and D’Hondt, S. (2016). Relationship of bacterial richness to organic degradation rate and sediment age in subseafloor sediment. *Appl. Environ. Microbiol.* 82, 4994–4999.
- Welhan, J.A., and Lupton, J.E. (1987). Light hydrocarbon gases in Guaymas basin hydrothermal fluids: thermogenic versus abiogenic origin. *AAPG Bull.* 71, 215–223.
- Willis, A., and Bunge, J. (2015). Estimating diversity via frequency ratios. *Biometrics* 71, 1042–1049.
- Xia, X., Guo, W., and Liu, H. (2017). Basin scale variation on the composition and diversity of archaea in the Pacific ocean. *Front. Microbiol.* 8, 2057.
- Yu, T., Wu, W., Liang, W., Lever, M.A., Hinrichs, K.U., and Wang, F. (2018). Growth of sedimentary Bathyarchaeota on lignin as an energy source. *Proc. Natl. Acad. Sci. U S A* 115, 6022–6027.
- Zhou, Z., Liu, Y., Lloyd, K.G., Pan, J., Yang, Y., Gu, J.D., and Li, M. (2019). Genomic and transcriptomic insights into the ecology and metabolism of benthic archaeal cosmopolitan, Thermopfundales (MBG-D archaea). *ISME J.* 13, 885–901.
- Ziervogel, K., and Arnosti, C. (2020). Substantial carbohydrate hydrolase activities in the water column of the Guaymas basin (Gulf of California). *Front. Mar. Sci.* 6, 815.

iScience, Volume 23

Supplemental Information

The Guaymas Basin Seafloor

Sedimentary Archaeome Reflects

Complex Environmental Histories

Gustavo A. Ramírez, Luke J. McKay, Matthew W. Fields, Andrew Buckley, Carlos Mortera, Christian Hensen, Ana Christina Ravelo, and Andreas P. Teske

1 *Supplemental Materials*

2 **The Guaymas Basin Subseafloor Sedimentary Archaeome Reflects Complex**
3 **Environmental Histories**

4
5 Gustavo A. Ramírez^{1,2*}, Luke J. McKay^{3,4}, Matthew W. Fields^{4,5}, Andrew Buckley¹, Carlos
6 Mortera⁶, Christian Hensen⁷, Ana Christina Ravelo⁸ & Andreas P. Teske¹

7
8 Author affiliations:

9 1: Department of Marine Sciences, University of North Carolina at Chapel Hill, NC, USA.

10 2: College of Veterinary Medicine, Western University of Health Sciences, Pomona, CA,
11 USA.

12 3: Department of Land Resources and Environmental Sciences, Montana State University,
13 Bozeman, MT, USA.

14 4: Center for Biofilm Engineering, Montana State University, Bozeman, MT, USA.

15 5: Department of Microbiology and Immunology, Montana State University, Bozeman, MT,
16 USA.

17 6: Instituto de Geofísica, Universidad Nacional Autónoma de México, Coyoacán, Mexico.

18 7: GEOMAR Helmholtz Centre for Ocean Research, Kiel, Germany.

19 8: Ocean Sciences Department, University of California, Santa Cruz, CA, USA.

20
21
22 *: Corresponding author

23
24
25 Correspondence:

26 Dr. Gustavo A. Ramírez

27 gara1985@email.unc.edu

28 gramirez@westernu.edu

29 zombiephylotype@gmail.com

30

31 **Transparent Methods**

32

33 *Sample Collection.* All samples were collected using piston coring during R/V *El Puma*
34 (Universidad Nacional Autónoma de México, UNAM) Expedition Guaymas14 to the Gulf of
35 California, October 14-27th, 2014. A 5-m long piston core (RNVP11) was obtained on Oct 21,
36 2014 from the central basin within the ring (27°N30.5090/111°W40.6860, 1749 m; core length
37 4.9 m), parallel to a control core (ContP10) approx. 1 mile to the west of Ringvent
38 (27°N30.5193/111°W42.1722; 1731 m depth, 3.93 m core length) collected on the same day.
39 Core SeepP06 was obtained on Oct. 19 from the lower Sonora Margin, near its boundary with
40 the Ridge flanks (27°N38.8367/111°W36.8595; 1681 m depth, 3.95 m core length). Core
41 OMZP12 was taken on Oct. 22 from the upper Sonora Margin (27°N52.1129/111°W41.5902,
42 667 m, 4 m core length) in the oxygen minimum zone as previously determined by water
43 column oxygen profiling (Calvert 1964). Core ContP03 was collected on Oct. 17 from the
44 northwestern end of the ridge flanks (27°N37.6759/ 111°W52.5740; 1611 m depth, 3.27 m
45 core length. Core ContP13 was obtained on Oct. 22 from the southeastern ridge flank of
46 Guaymas Basin (27°N12.4470/111°W13.7735, 1859m depth, 3.31 m core length).

47

48 *Geochemical Analyses.* Porewater was obtained from freshly collected sediments on RV *El*
49 *Puma* by centrifuging ca. 40 ml sediment samples in 50 ml conical Falcon tubes for ca. 5 to 10
50 minutes, using a Centra CL-2 Tabletop centrifuge (Thermo Scientific) at 1000g, until the
51 sediment had settled and produced ca. 8 to 10 ml of porewater. Porewater was extracted from
52 5 cm thick sediment samples, which are designated by the top of each sample. For example, a
53 “95 cm” geochemistry sample extends from 95 to 100 cm below the sediment surface. Filtered,
54 unamended, porewater samples prepared shipboard were stored at 4°C for shored-based
55 analyses. Sulfate, sulfide, methane, and DIC porewater profiles for cores SeepP06, ContP10,
56 RNVP11, and OMZP12 were previously published (Teske et al 2019), and are re-plotted here
57 for comparison with unpublished profiles from cores ContP03 and ContP13. Porewater
58 analyses were performed as previously described, using the colorimetric Cline assay for
59 sulfide, ion chromatography for sulfate, and GC-IRMS for DIC and methane (Teske et al
60 2019). Carbon and nitrogen isotopic and elemental composition was determined at the Stable
61 Isotope Laboratory (SIL) at the University of California, Santa Cruz (UCSC). Bulk sediment
62 $\delta^{15}\text{N}$ and elemental ratio data were collected using 20 mg samples in Sn capsules; organic $\delta^{13}\text{C}$
63 and elemental composition data were collected using 2.5 mg samples of acidified sediment in

64 Sn capsules. All samples were measured by Dumas combustion performed on a Carlo Erba
65 1108 elemental analyzer coupled to a ThermoFinnigan Delt Plus XP isotope ratio mass
66 spectrometer (EA-IRMS). An in-house gelatin standard, Acetanilide, and an in-house bulk
67 sediment standard, “Monterey Bay Sediment Standard”, were used in all runs. Reproducibility
68 of an in-house matrix-matched sediment standard is <0.1‰ VPDB for $\delta^{13}\text{C}$ and <0.2‰ AIR
69 for $\delta^{15}\text{N}$. Data is corrected for blank, and for drift when appropriate. Carbon and nitrogen
70 elemental composition was estimated based on standards of known composition, for which
71 analytical precision is determined to be better than 1 %. Filtered but unamended porewater
72 samples, stored at 4°C, were used for quantifying multiple stable ions, including silicate, by
73 ion chromatography at GEOMAR, Kiel, Germany (Hensen et al 2007). All geochemical data
74 in this study are publicly available at the Biological and Chemical Oceanography Data
75 Management Office (BCO-DMO) under the following dataset IDs: 661750, 661658, 66175
76 and 661808 for methane, DIC, sulfate and sulfide, respectively.

77

78 *3. DNA extraction and gene sequencing*

79 Samples for DNA sequencing [approx. 2 cm³ each] were obtained by syringe coring at the
80 indicated depth [in cm] below the sediment surface. Freshly collected samples were
81 immediately frozen (-80°C) for storage and transport back to shore. DNA for all survey sites
82 was extracted from ~0.5-1.0 cm³ sediment sample volumes using the Powersoil DNA
83 extraction kit according to the manufacturer’s instructions (QIAGEN, Carlsbad, CA, USA).
84 Archaeal 16S rRNA gene amplicons from DNA extracts were generated using the following
85 primer set: A751F: 5’-CGA CGG TGA GRG RYG AA-3’ and A1204R: 5’-TTM GGG GCA
86 TRC NKA CCT-3’ using the following thermocycling program: initial denaturation for 2 mins
87 at 94°C, 30 x [94°C for 1 min, 55°C for 1 min, 72°C for 1 min], and a final 10 min extension
88 at 72°C, as suggested elsewhere (Baker et al 2003). Amplicons were sequenced on an Illumina
89 MiSeq platform (Illumina, San Diego, CA, USA) at the Center for Biofilm Engineering in
90 Bozeman, Montana. Sequencing run specifications are found in the Visualization and Analysis
91 of Microbial Population Structures (VAMPSs) website
92 (<https://vamps.mbl.edu/resources/primers.php>) (Huse et al 2014).

93

94 *4. Sequence Processing*

95 Sequences were processed with *mothur* v.1.39.5 (Schloss et al 2009) following the *mothur*
96 Illumina MiSeq SOP (Kozich et al 2013). Briefly, forward and reverse reads were merged into

97 contigs and selected based on primer-specific amplicon length and the following parameters:
98 maximum homopolymers of 6bp, and zero ambiguities. High quality sequences were aligned
99 against the *mothur*-recreated Silva SEED v132 database (Yarza et al 2010) and subsequently
100 pre-clustered at 1% dissimilarity. As suggested elsewhere (Kozich et al 2013), spurious
101 sequences are mitigated by abundance ranking and merging with rare sequences based on
102 minimum differences of three base pairs. Chimeras were detected and removed using UCHIME
103 de novo mode (Edgar et al 2011). Sequences were then clustered, by generating a distance
104 matrix using the average neighbor method, into operational taxonomic units (OTUs, 97%
105 similarity cutoff). OTU classification was performed on *mothur* using the SILVA v132
106 database as implemented using the classify.seqs command using the Wang algorithm (kmer
107 assignment with 1/8 kmer replacement as bootstrap) and cutoff=80 (minimal bootstrap value
108 for sequence taxonomy assignment). All sequence data are publically available at the following
109 repository: NCBI under BioProject PRJNA553578 and accession numbers SRX6444849-
110 SRX6444877.

111

112 5. Sequence Analyses

113 5.1 Community Analyses and Visualizations

114 Community analyses were performed in *RStudio* version 0.98.1091 (Racine 2012),
115 implemented in R version 3.5.2, using the *vegan* (Oksanen et al 2015) and *phyloseq* (McMurdie
116 and Holmes 2013) R-packages. Sample richness analyses used the R package *breakaway*
117 (Willis et al. 2017) for inferring precision of diversity estimations given the heterologous
118 sequencing depth. Data were rlog normalized using *DESeq2* (Love et al 2014) prior to
119 ordination using Bray-Curtis distances. An identical normalization strategy was used on Bray-
120 Curtis distances for co-occurrence network analysis performed using the *makenetwork()*
121 *phyloseq* command and visualized using the *igraph* R-package. *DESeq2* was also used to
122 perform differential abundance analyses of taxa with low abundance taxa ($n < 100$ total reads
123 per OTU) removed for the un-rarefied dataset, as suggested elsewhere (McMurdie and Holmes
124 2014).

125

126 5.2 Phylogenetic Analyses

127 Sequence alignments were performed using the high speed multiple sequence alignment
128 program MAFFT (Katoh and Standley 2013) with the command: `mafft --maxiterate 1000 --
129 localpair seqs.fasta > aligned.seqs.fasta`. Maximum likelihood trees with 100 bootstrap support
130 were constructed using the RAxML (Stamatakis 2014) program using the following

131 parameters: raxmlHPC -f a -m GTRGAMMA -p 12345 -x 12345 -# 100 -s aligned.seqs.fasta -
132 n T.tree, -T 4 ML search + bootstrapping. Newick trees files were uploaded to FigTree v1.4.2
133 for visualization.

<i>Core ID</i>	Latitude	Longitude	Collection Date (2014)	Core Length (m)	Water Depth (m)
<i>ContP3</i>	27°N37.6759	111°W52.5740	Oct. 17	3.27	1611
<i>SeepP6</i>	27°N38.8367	111°W36.8595	Oct. 19	3.95	1681
<i>ContP10</i>	27°N30.5193	111°W42.1722	Oct. 21	3.93	1731
<i>RNVP11</i>	27°N30.5090	111°W40.6860	Oct. 21	4.9	1749
<i>OMZP12</i>	27°N52.1129	111°W41.5902	Oct. 22	4	667
<i>ContP13</i>	27°N12.4470	111°W13.7735	Oct. 22	3.31	1859

134
135
136

Table S1. Related to Figure 1. Core site metadata.

Sample	Lokiarchaea
ContP03_9	0.019%
ContP03_104	0.007%
ContP03_202	0.000%
ContP03_301	0.000%
SeepP06_5	0.024%
SeepP06_105	0.009%
SeepP06_205	0.000%
SeepP06_304	0.000%
SeepP06_394	0.000%
ContP10_5	0.012%
ContP10_104	0.000%
ContP10_204	0.000%
ContP10_303	0.000%
ContP10_378	0.000%
RNVP11_5	0.000%
RNVP11_95	0.020%
RNVP11_195	0.000%
RNVP11_295	0.000%
RNVP11_394	0.000%
RNVP11_486	0.000%
OMZP12_5	0.003%
OMZP12_105	0.000%
OMZP12_204	0.000%
OMZP12_304	0.000%
OMZP12_379	0.003%
ContP13_5	0.006%
ContP13_111	0.002%
ContP13_210	0.000%
ContP13_310	0.000%

137

138

139

140

Table S2. Related to Figure 6. Percent of total community contribution of Lokiarchaea sequences in all samples based on SILVA132 taxonomic assignments.

Core_cmbsf	All_ANME	ANME-2		
		ANME-1	ANME-2a-2b	ANME-2c
ContP3_009	0.034	0.000	0.000	0.034
ContP3_104	0.002	0.002	0.000	0.002
ContP3_202	0.000	0.000	0.000	0.000
ContP3_301	0.000	0.000	0.000	0.000
SeepP6_005	0.030	0.018	0.012	0.030
SeepP6_105	8.863	8.863	0.000	8.863
SeepP6_205	32.063	32.063	0.000	32.063
SeepP6_304	32.446	32.440	0.006	32.446
SeepP6_394	39.810	39.810	0.000	39.810
ContP10_005	0.111	0.088	0.024	0.111
ContP10_104	0.092	0.092	0.000	0.092
ContP10_204	0.003	0.000	0.003	0.003
ContP10_303	0.447	0.447	0.000	0.447
ContP10_378	0.000	0.000	0.000	0.000
RNVP11_005	0.009	0.009	0.000	0.009
RNVP11_095	0.000	0.000	0.000	0.000
RNVP11_195	0.988	0.988	0.000	0.988
RNVP11_295	0.000	0.000	0.000	0.000
RNVP11_394	0.000	0.000	0.000	0.000
RNVP11_486	0.000	0.000	0.000	0.000
OMZP12_005	0.000	0.000	0.000	0.000
OMZP12_105	0.123	0.121	0.002	0.123
OMZP12_204	2.098	2.098	0.000	2.098
OMZP12_304	0.629	0.629	0.000	0.629
OMZP12_379	0.967	0.967	0.000	0.967
ContP13_005	0.476	0.429	0.029	0.476
ContP13_111	0.006	0.002	0.004	0.006
ContP13_211	0.055	0.012	0.043	0.055
ContP13_310	0.004	0.000	0.004	0.004

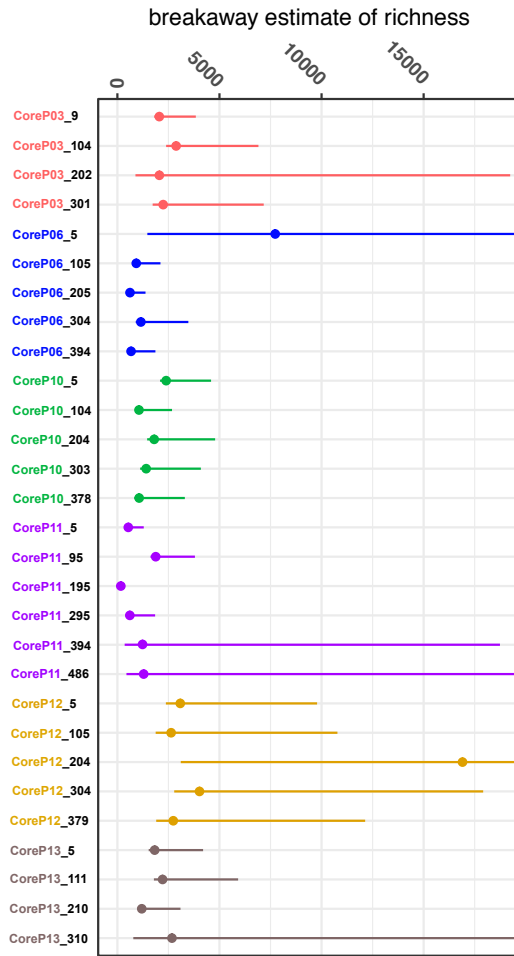
178
179
180
181
182
183
184
185
186
187
188
189

Table S3. Related to Figure 6. Percent of total community contribution of ANME sequences in all samples based on SILVA132 taxonomic assignments. The All_ANME column shows the percent contribution of sequences classified as ANME in each sample. Columns ANME-1, ANME-2a-2b, and ANME-2c show the percent breakdown of the respective ANME lineages in each sample and their sum is equal to the All_ANME column percentage.

<u>Sample Name</u>	<u>DNA yield (ng/μL)</u>	<u>Num. of seqs post Mothur QC and chimera removal</u>
ContP3_9	7	21,443
ContP3_104	6.9	47,239
ContP3_202	6.6	16,038
ContP3_301	9.4	45,559
SeepP6_5	9	17,196
SeepP6_105	4.3	11,595
SeepP6_205	9.1	9,274
SeepP6_304	9.4	18,043
SeepP6_394	8	10,047
ContP10_5	9.2	25,975
ContP10_104	7.7	12,289
ContP10_204	8	35,076
ContP10_303	14.5	29,782
ContP10_378	7.6	25,682
RNVP11_5	6.7	11,184
RNVP11_95	6.7	30,452
RNVP11_195	7.1	2,978
RNVP11_295	7	19,515
RNVP11_394	7.4	14,142
RNVP11_468	7.9	29,851
OMZP12_5	7.9	63,690
OMZP12_105	9	51,384
OMZP12_204	7.8	167,234
OMZP12_304	7.3	154,763
OMZP12_379	8.1	76,729
ContP13_5	6.6	17,573
ContP13_111	7.9	47,432
ContP13_210	6.8	25,989
ContP13_310	7.3	24,873

190
191
192

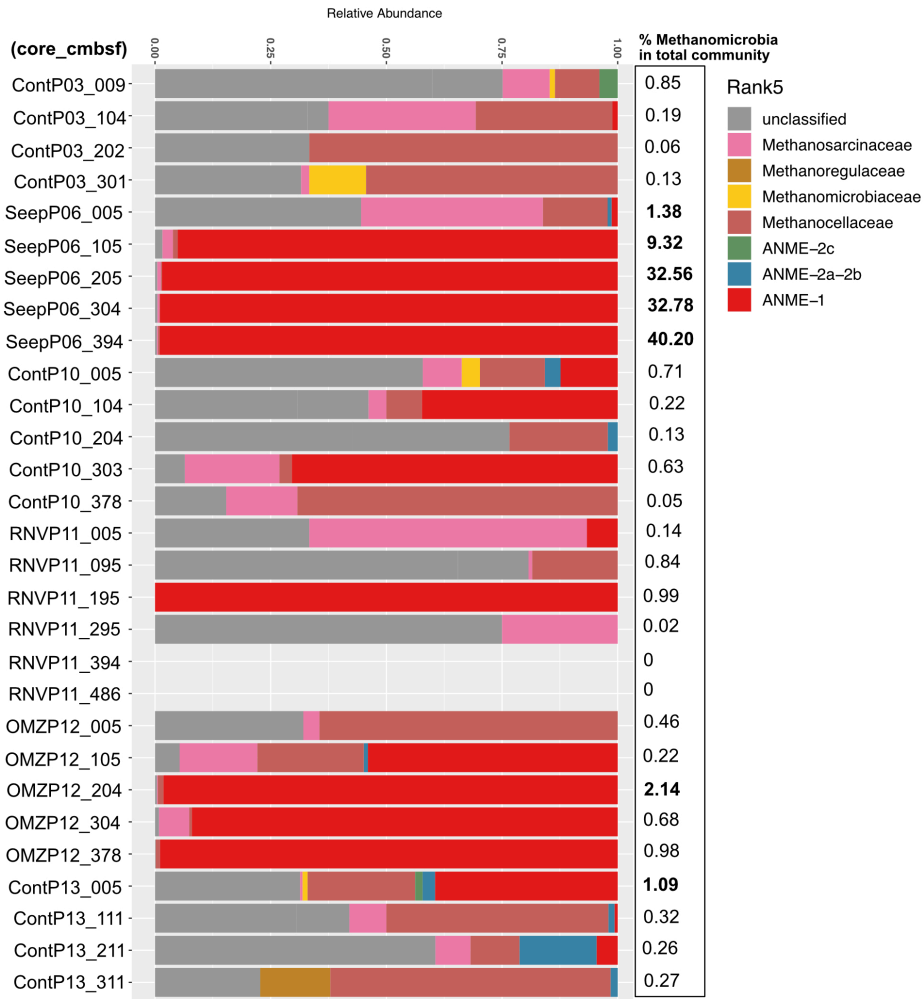
Table S4. Related to Figure 3. Total DNA yield and high-quality sequence numbers for all samples.



193
 194
 195
 196
 197

Figure S1. Related to Figure 3. Breakaway estimate of total species richness with model confidence intervals for color-coded cored site for all depths.

Guaymas Basin Methanomicrobia Community Composition (SILVA 132 Rank5)



198
199
200
201
202

Figure S2. Related to Figure 6. Methanomicrobia community composition for all cores in this survey.

203
204
205
206
207
208
209
210
211
212
213
214
215
216
217
218
219
220
221
222
223
224
225
226
227
228
229
230
231
232
233
234
235
236
237
238
239
240
241
242
243
244
245
246
247
248
249
250
251
252

Works Cited

- Baker GC, Smith JJ, Cowan DA (2003). Review and re-analysis of domain-specific 16S primers. *J of Microbiol Meth* **55**: 541-555.
- Calvert SE (1964). "Factors affecting distribution of laminated diatomaceous sediments in the Gulf of California" In *Marine Geology of the Gulf of California*, Vol. 3, eds. T.H. van Andel and G.G. Shor, (Tulsa: American Association of Petroleum Geologists Memorir) **3**: 311-330.
- Edgar RC, Haas BJ, Clemente JC, Quince C, Knight R (2011). UCHIME improves sensitivity and speed of chimera detection. *Bioinformatics* **27**: 2194-2200.
- Hensen C, Nuzzo M, Hornibrook E, Pinheiro LM, Bock B, Magalhães VH *et al* (2007). Sources of mud volcano fluids in the Gulf of Cadiz—indications for hydrothermal imprint. *Geochim et Cosmochim Acta* **71**: 1232-1248.
- Huse SM, Mark Welch D, Voorhis A, Shipunova A, Morrison HG, Eren AM *et al* (2014). VAMPS: a website for visualization and analysis of microbial population structures. *BMC Bioinformatics* **15**.
- Katoh K, Standley DM (2013). MAFFT multiple sequence alignment software version 7: improvements in performance and usability. *Mol Biol Evol* **30**: 772-780.
- Kozich JJ, Westcott SL, Baxter NT, Highlander SK, Schloss PD (2013). Development of a dual-index sequencing strategy and curation pipeline for analyzing amplicon sequence data on the MiSeq Illumina sequencing platform. *Appl Environ Microbiol* **79**: 5112-5120.
- Love MI, Huber W, Anders S (2014). Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biol* **15**: 550.
- McMurdie PJ, Holmes S (2013). phyloseq: an R package for reproducible interactive analysis and graphics of microbiome census data. *PLoS One* **8**: e61217.
- McMurdie PJ, Holmes S (2014). Waste not, want not: why rarefying microbiome data is inadmissible. *PLoS Comput Biol* **10**.
- Oksanen J, Blanchet FG, Kindt R, Legendre P, Minchin PR, O'Hara RB *et al* (2015). vegan: Community Ecology Package. R Package Version 2.2-1. Available online at: <http://CRANR-project.org/package=vegan>.
- Racine JS (2012). RStudio: A Platform-Independent IDE for R and Sweave. *Journal of Applied Econometrics* **27**: 167-172.
- Schloss PD, Westcott SL, Ryabin T, Hall JR, Hartmann M, Hollister EB *et al* (2009). Introducing mothur: open-source, platform-independent, community-supported software for describing and comparing microbial communities. *Appl Environ Microbiol* **75**: 7537-7541.
- Stamatakis A (2014). RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics* **30**: 1312-1313.

253 Teske A, McKay L, Ravelo AC, Aiello I, Mortera C, Núñez-Useche F *et al* (2019).
254 Characteristics and evolution of sill-driven off-axis hydrothermalism in Guaymas Basin- the
255 Ringvent site. *Scientific Reports* **9**.
256
257 Yarza P, Ludwig W, Euzeby J, Amann R, Schleifer KH, Glöckner FO *et al* (2010). Update of
258 the All-Species Living Tree Project based on 16S and 23S rRNA sequence analyses. *Syst Appl*
259 *Microbiol* **33**: 291-299.
260
261