Supplemental Information

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Sedimentary Archaeome Reflects

Complex Environmental Histories

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Supplemental Materials

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**Transparent Methods**

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

**Geochemical Analyses.** Porewater was obtained from freshly collected sediments on RV *El Puma* by centrifuging ca. 40 ml sediment samples in 50 ml conical Falcon tubes for ca. 5 to 10 minutes, using a Centra CL-2 tabletop centrifuge (Thermo Scientific) at 1000 g, until the sediment had settled and produced ca. 8 to 10 ml of porewater. Porewater was extracted from 5 cm thick sediment samples, which are designated by the top of each sample. For example, a “95 cm” geochemistry sample extends from 95 to 100 cm below the sediment surface. Filtered, unamended, porewater samples prepared shipboard were stored at 4°C for shore-based analyses. Sulfate, sulfide, methane, and DIC porewater profiles for cores SeepP06, ContP10, RNVP11, and OMZP12 were previously published (Teske et al 2019), and are re-plotted here for comparison with unpublished profiles from cores ContP03 and ContP13. Porewater analyses were performed as previously described, using the colorimetric Cline assay for sulfide, ion chromatography for sulfate, and GC-IRMS for DIC and methane (Teske et al 2019). Carbon and nitrogen isotopic and elemental composition was determined at the Stable Isotope Laboratory (SIL) at the University of California, Santa Cruz (UCSC). Bulk sediment δ15N and elemental ratio data were collected using 20 mg samples in Sn capsules; organic δ13C and elemental composition data were collected using 2.5 mg samples of acidified sediment in
Sn capsules. All samples were measured by Dumas combustion performed on a Carlo Erba 1108 elemental analyzer coupled to a ThermoFinnigan Delt Plus XP isotope ratio mass spectrometer (EA-IRMS). An in-house gelatin standard, Acetanilide, and an in-house bulk sediment standard, “Monterey Bay Sediment Standard”, were used in all runs. Reproducibility of an in-house matrix-matched sediment standard is <0.1‰ VPDB for δ^{13}C and <0.2‰ AIR for δ^{15}N. Data is corrected for blank, and for drift when appropriate. Carbon and nitrogen elemental composition was estimated based on standards of known composition, for which analytical precision is determined to be better than 1%. Filtered but unamended porewater samples, stored at 4°C, were used for quantifying multiple stable ions, including silicate, by ion chromatography at GEOMAR, Kiel, Germany (Hensen et al 2007). All geochemical data in this study are publicly available at the Biological and Chemical Oceanography Data Management Office (BCO-DMO) under the following dataset IDs: 661750, 661658, 66175 and 661808 for methane, DIC, sulfate and sulfide, respectively.

3. DNA extraction and gene sequencing

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

4. Sequence Processing

Sequences were processed with mothur v.1.39.5 (Schloss et al 2009) following the mothur Illumina MiSeq SOP (Kozich et al 2013). Briefly, forward and reverse reads were merged into
contigs and selected based on primer-specific amplicon length and the following parameters: maximum homopolymers of 6bp, and zero ambiguities. High quality sequences were aligned against the mothur-recreated Silva SEED v132 database (Yarza et al 2010) and subsequently pre-clustered at 1% dissimilarity. As suggested elsewhere (Kozich et al 2013), spurious sequences are mitigated by abundance ranking and merging with rare sequences based on minimum differences of three base pairs. Chimeras were detected and removed using UCHIME de novo mode (Edgar et al 2011). Sequences were then clustered, by generating a distance matrix using the average neighbor method, into operational taxonomic units (OTUs, 97% similarity cutoff). OTU classification was performed on mothur using the SILVA v132 database as implemented using the classify.seqs command using the Wang algorithm (kmer assignment with 1/8 kmer replacement as bootstrap) and cutoff=80 (minimal bootstrap value for sequence taxonomy assignment). All sequence data are publically available at the following repository: NCBI under BioProject PRJNA553578 and accession numbers SRX6444849-SRX6444877.

5. Sequence Analyses

5.1 Community Analyses and Visualizations

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

5.2 Phylogenetic Analyses

Sequence alignments were performed using the high speed multiple sequence alignment program MAFFT (Katoh and Standley 2013) with the command: mafft --maxiterate 1000 --localpair seqs.fasta > aligned.seqs.fasta. Maximum likelihood trees with 100 bootstrap support were constructed using the RAxML (Stamatakis 2014) program using the following
parameters: raxmlHPC -f a -m GTRGAMMA -p 12345 -x 12345 -# 100 -s aligned.seqs.fasta -n T.tree, -T 4 ML search + bootstrapping. Newick trees files were uploaded to FigTree v1.4.2 for visualization.
| Core ID | Latitude    | Longitude    | Collection Date (2014) | Core Length (m) | Water Depth (m) |
|---------|-------------|--------------|------------------------|-----------------|-----------------|
| ContP3  | 27°N37.6759 | 111°W52.5740 | Oct. 17                | 3.27            | 1611            |
| SeepP6  | 27°N38.8367 | 111°W36.8595 | Oct. 19                | 3.95            | 1681            |
| ContP10 | 27°N30.5193 | 111°W42.1722 | Oct. 21                | 3.93            | 1731            |
| RNVP11  | 27°N30.5090 | 111°W40.6860 | Oct. 21                | 4.9             | 1749            |
| OMZP12  | 27°N52.1129 | 111°W41.5902 | Oct. 22                | 4               | 667             |
| ContP13 | 27°N12.4470 | 111°W13.7735 | Oct. 22                | 3.31            | 1859            |

Table S1. Related to Figure 1. Core site metadata.
Table S2. Related to Figure 6. Percent of total community contribution of Lokiarchaea sequences in all samples based on SILVA132 taxonomic assignments.
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.

| Core.cmbsf      | All_ANME | ANME-1 | ANME-2a-2b | ANME-2c |
|-----------------|----------|--------|------------|--------|
| ContP3_009      | 0.034    | 0.000  | 0.000      | 0.004  |
| ContP3_104      | 0.002    | 0.002  | 0.000      | 0.000  |
| 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      | 0.000  |
| SeepP6_105      | 8.863    | 8.863  | 0.000      | 0.000  |
| SeepP6_205      | 32.063   | 32.063 | 0.000      | 0.000  |
| SeepP6_304      | 32.446   | 32.440 | 0.006      | 0.000  |
| SeepP6_394      | 39.810   | 39.810 | 0.000      | 0.000  |
| ContP10_005     | 0.111    | 0.088  | 0.024      | 0.000  |
| ContP10_104     | 0.092    | 0.092  | 0.000      | 0.000  |
| ContP10_204     | 0.003    | 0.000  | 0.003      | 0.000  |
| ContP10_303     | 0.447    | 0.447  | 0.000      | 0.000  |
| ContP10_378     | 0.000    | 0.000  | 0.000      | 0.000  |
| RNVP11_005      | 0.009    | 0.009  | 0.000      | 0.000  |
| RNVP11_095      | 0.000    | 0.000  | 0.000      | 0.000  |
| RNVP11_195      | 0.988    | 0.988  | 0.000      | 0.000  |
| 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      | 0.000  |
| OMZP12_204      | 2.098    | 2.098  | 0.000      | 0.000  |
| OMZP12_304      | 0.629    | 0.629  | 0.000      | 0.000  |
| OMZP12_379      | 0.967    | 0.967  | 0.000      | 0.000  |
| ContP13_005     | 0.476    | 0.429  | 0.029      | 0.000  |
| ContP13_111     | 0.006    | 0.002  | 0.004      | 0.000  |
| ContP13_211     | 0.055    | 0.012  | 0.043      | 0.000  |
| ContP13_310     | 0.004    | 0.000  | 0.004      | 0.000  |
| Sample Name  | DNA yield (ng/µL) | Num. of seqs post Mothur QC and chimera removal |
|--------------|-------------------|-----------------------------------------------|
| 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                                        |

Table S4. Related to Figure 3. Total DNA yield and high-quality sequence numbers for all samples.
Figure S1. Related to Figure 3. Breakaway estimate of total species richness with model confidence intervals for color-coded cored site for all depths.
Figure S2. Related to Figure 6. Methanomicrobia community composition for all cores in this survey.
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