Genomic insights into potential interdependencies in microbial hydrocarbon and nutrient cycling in hydrothermal sediments

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Abstract

Background: Deep-sea hydrothermal vents are hotspots for productivity and biodiversity. Thermal pyrolysis and circulation produce fluids rich in hydrocarbons and reduced compounds that stimulate microbial activity in surrounding sediments. Several studies have characterized the diversity of Guaymas Basin (Gulf of California) sediment-inhabiting microorganisms; however, many of the identified taxa lack cultures or genomic representations. Here, we resolved the metabolic potential and community-level interactions of these diverse communities by reconstructing and analyzing microbial genomes from metagenomic sequencing data.

Results: We reconstructed 115 microbial metagenome-assembled genomes comprising 27 distinct archaeal and bacterial phyla. The archaea included members of the DPANN and TACK superphyla, Bathyarchaeota, novel Methanosarcinales (GoM-Arc1), and anaerobic methane-oxidizing lineages (ANME-1). Among the bacterial phyla, members of the Bacteroidetes, Chloroflexi, and Deltaproteobacteria were metabolically versatile and harbored potential pathways for hydrocarbon and lipid degradation and a variety of respiratory processes. Genes encoding enzymes that activate anaerobic hydrocarbons for degradation were detected in Bacteroidetes, Chloroflexi, Latescibacteria, and KSB1 phyla, while the reconstructed genomes for most candidate bacteria phyla (Aminicenantes, Atribacteria, Omnitrophica, and Stahlbacteria) indicated a fermentative metabolism. Newly obtained GoM-Arc1 archaeal genomes encoded novel pathways for short-chain hydrocarbon oxidation by alkyl-coenzyme M formation. We propose metabolic linkages among different functional groups, such as fermentative community members sharing substrate-level interdependencies with sulfur- and nitrogen-cycling microbes.

Conclusions: Overall, inferring the physiologies of archaea and bacteria from metagenome-assembled genomes in hydrothermal deep-sea sediments has revealed potential mechanisms of carbon cycling in deep-sea sediments. Our results further suggest a network of biogeochemical interdependencies in organic matter utilization, hydrocarbon degradation, and respiratory sulfur cycling among deep-sea-inhabiting microbial communities.

Keywords: Deep sea, Microbiome, Metagenome, Bacteria, Archaea, Population genomes, Comparative genomics

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Background

Marine sediments form the largest repository of organic carbon and the most extensive habitat for microbial life, where sediment-inhabiting microorganisms drive nutrient and carbon cycling [1–4]. Guaymas Basin (GB), a hydrothermally active seafloor-spraying center in the Gulf of California, is characterized by high primary production, rapid sedimentation and deposition, and hydrothermal processing of buried organic matter within its massive sediment cover [5, 6]. Hydrothermal alterations transform the deposited carbon and produce large amounts of methane, petroleum-like compounds (alkanes and polycyclic aromatic hydrocarbons (PAHs)), organic acids, and ammonia [5, 7–9]. These substrates get distributed throughout the sediments by hydrothermal circulation and are readily assimilated by the local microbial community [10]. For example, hydrothermal fluids containing high concentrations of methane (> 15 mM) mixing with seawater sulfate (28 mM) favor the anaerobic oxidation of methane (AOM), which occurs at high temperatures in Guaymas Basin sediments [11–13]. AOM is usually carried out in a syntrophic relationship between anaerobic methanotrophic (ANME) archaea and members of the Deltaproteobacteria that couple the oxidation of methane with the reduction of sulfate [14]. In hydrothermal sediments of Guaymas Basin, AOM is also performed at high temperatures by a syntrophic consortium of ANME-1 archaea and the deep-branched, hydrogenotrophic sulfate-reducing bacterium Candidatus Desulfovviridus auxilli [15–17]. In contrast to this well-studied interaction, little is known about the degradation of other abundant hydrocarbons (such as PAHs and alkanes) by the GB microbiome.

The microbial community composition of GB sediments has been thoroughly described using marker-gene studies [12, 13]. For example, an automated ribosomal spacer analysis indicated that > 80% of the detected operational taxonomic units (OTUs) were shared across different temperature and depth profiles, suggesting a high connectivity across sediments [12, 18]. Furthermore, depth and temperature regime are thought to influence community assembly [18]. Temperatures in GB sediments range from ~ 3 °C at the surface up to ~ 200 °C at 30- to 50-cm depth, which is accompanied with geochemical zonation due to mixing of hot vent fluids with cold ocean water [12, 18, 19]. 16S rRNA gene sequencing of Guaymas Basin sediments has revealed numerous bacterial and archaeal lineages [12, 13]. Among the archaea, ANME-1 members are frequently detected, as well as other common deep-sea lineages including Marine Benthic Group D (MBG-D) and Bathyarchaeota (formerly MCG). Consistently detected bacterial community members include Epsilonproteobacteria, Deltaproteobacteria, as well as the uncultured SEEP-SRB2, Candidatus Desulfovviridus auxilli (previously HotSeep-1) lineages, Bacteroidetes, or Chloroflexi [12, 13]. Despite our knowledge about the geochemistry of GB sediments, and the taxonomic composition of its microbial communities, single-gene studies lack insights into the metabolic capacities and ecological connectivity in this unique deep-sea environment. To address this gap, we obtained metagenomic libraries from two GB sediment sites. Accompanying biogeochemical data for these two sites suggest that in addition to abundant methane, short-chain alkanes (C1 to C5) are available in substantial concentrations (20 to 100 μM) [13]. Consistent with the frequent detection of sulfate-reducing microbial populations that use hydrogen as preferred electron donor, porewater hydrogen concentrations are consistently low (< 10 nM) [20]. Therefore, we hypothesize that this habitat is suitable for syntrophic alkane oxidation by consortia of alkane-oxidizing archaea and sulfate-reducing bacteria that catalyze the terminal electron transfer from alkane to sulfate. Additionally, we aim to address whether the deposition and subsequent alteration of abundant organic carbon favors an unusually diverse microbial community in Guaymas Basin sediments [5, 6]. From the two GB sediment sites, we were able to reconstruct 115 metagenome-assembled genomes (MAGs) belonging to 27 bacterial and archaeal phyla, allowing us to begin to resolve the metabolic potential of a variety of uncultured community members. Physiological analyses of these communities revealed novel pathways for hydrocarbon processing and potential ecological interdependencies.

Results

Genome reconstructions and community composition

We obtained ~ 242 gigabases of Illumina shotgun sequencing data from two GB hydrothermal sediment sites, where sediments cores were collected during Alvin dives 4484 and 4572 in December 2008 and 2009, respectively (Additional file 1: Tables S1 and S2, details in the “Methods” section). Samples from two depth profiles from each of these two representative sites were selected for sequencing based on the availability of accompanying biogeochemical data, thermal gradients, and evidence of a diverse microbial community [12, 13, 21]. De novo assembly and tetranucleotide (and coverage) binning of sequencing data from these four samples allowed the reconstruction of 38 archaeal and 77 bacterial draft metagenome-assembled genomes (MAGs, completeness > 50%, Additional file 1: Table S3, Figure S1). Of those MAGs, 51 were estimated to be > 70% and 20 to be > 80% complete, with minimal single-gene duplications (≤ 10%, Additional file 1: Table S3).

The reconstructed MAGs comprise a total of 9 archaeal and 18 bacterial phyla (Figs. 1 and 2, Additional file 1: Figures S2–S7, Table S3). Overall, this community is taxonomically diverse and includes lineages of the Bathyarchaeota, Methanosarcinales (novel GoM-Arc1), Methanomicrobia (ANME-1 groups), Thermoplastales
(uncultured VC2.1 Arc6, CCA47 lineages), Bacteroidetes, Chloroflexi, Deltaproteobacteria, and Gammaproteobacteria. Additionally, we identified several MAGs from a variety of little understood candidate phyla: Aminicenantes (OP8), Atribacteria (OP9), Cloacimonas, Omnitrophica (WOR-2), Latescibacteria (WS3), WOR-3, Zixibacteria, as well as Pacearchaeota and Geothermarchaeota. Phylum WOR-3 MAGs were first recovered from estuary sediments and have also been seen in groundwater sediments [22, 23]; we propose they be named Candidatus “Stahlbacteria” after Dr. David Stahl, an accomplished environmental microbiologist and early proponent of 16S rRNA-based
microbial ecology. The placement of Candidatus Stahlbacter as a new phylum was supported both by a phylogenetic analysis using 16 concatenated ribosomal marker genes as well as an extended marker set using 37 concatenated genes for a more robust phylogenetic placement (see the “Methods” section; Fig. 2, Additional file 1: Figure S7). The most abundant MAGs at both sites were assigned to a gammproteobacterium of the family Beggiatoaceae (ex4572_84) and an archaeum of the phylum Euryarchaeota of the class Methanomicrobia (ex4572_4) (Additional file 1: Figures S8, S9). We did detect a few site-specific differences, such as the presence of Omnitrophica in samples from dive 4484 but not 4572, suggesting there are community-level differences across sample locations and/or years (Additional file 1: Figures S8, S9). However, to further resolve potential site-/time-
dependent genome-level differences, a broader sampling scale would be required.

**Carbon metabolism**

To infer the metabolic potential of GB community members, we annotated genes within each of the MAGs using a variety of protein databases (see the “Methods” section). Functional interpretations were aided by linking individual genes to complete pathways within each MAG. Notably, the basic features of inferred physiologies were fairly consistent within organisms of the same phyla, allowing for MAGs to be grouped for metabolic comparisons (Figs. 1 and 2, Additional file 1: Table S4).

Pathways involved in the degradation of detrital organic matter (including complex carbohydrates, lipids, and proteins) were prevalent across both archaeal and bacterial MAGs (Fig. 3). The most numerous carbohydrate-degradation genes encoded for alpha-amylase (starch degradation), members of the glycoside hydrolase family 3 (GH3; cellulose degradation), and endoglucanase (cellulose degradation). Several carbohydrate-degradation genes that were abundant in bacteria, including GH3, beta-D-glucoronidase, pullulanase, and beta-1,4-mannosidase, were rare or even absent from archaea (Additional file 1: Table S5, \( P \) value < 0.05 based on a non-parametric Mann–Whitney test, Bonferroni corrected). This finding suggests that bacteria have access to a more diverse carbohydrate pool. *Chloroflexi*, KS1, *Spirochaetes*, *Methanosarcinales*, and *Thermofilum* contained the highest number of carbohydrate-degrading genes, while no clear lineage-specific pattern was observed for peptidases (Fig. 4, Additional file 1: Figure S10). Aside from carbohydrate-degradation, the ability to degrade lipids via the beta-oxidation pathway was common to members of *Bacteroidetes*, *Chloroflexi*, and *Deltaproteobacteria*. Additionally, we identified the key beta-oxidation genes for acyl-CoA dehydrogenase, enoyl-CoA hydratase, 3-hydroxyacyl-CoA dehydrogenase, and acetyl-CoA C-
acyltransferase in Geothermarchaeota, Archaeoglobales, and the majority of the Bathyarchaeota genomes.

The ability to ferment (via glycolysis) appeared to be almost universal among GB archaea and bacteria (Figs. 1 and 2, Additional file 1: Table S4). Bacteria likely ferment various carbon sources, including complex carbohydrates (cellulose, hydrocarbons) and peptides (after hydrolysis into monomeric sugars and amino acids) predominantly into hydrogen and ethanol, followed by lactate and acetate. Fe,Fe-hydrogenases for producing H₂ were identified in Bacteroidetes, Fusobacteria, and Omnitrophica [24]. The archaea identified are primarily predicted to produce hydrogen and acetate, and none appear to be able to produce lactate. Among the archaea, Ni,Fe-hydrogenases were found in the majority of phyla and orders, including Aenigmarchaeota, Desulfurococcales, and Thermoplasmales, while Fe,Fe-hydrogenases were found in various bins but could not be consistently linked to specific lineages.

Hydrocarbon utilization

Short-chain (petroleum-like) C₂-C₁₀ hydrocarbons are rapidly generated by hydrothermal pyrolysis and thus are a particularly abundant source of carbon and energy in hot GB sediments [5]. No genes for the aerobic degradation of these hydrocarbons were identified in the GB MAGs, but genes for anaerobic hydrocarbon degradation were detected among several phyla, including Bacteroidetes, Chloroflexi, Latescibacteria, and KSB1 contain genes that potentially encode for the benzyllsuccinate synthase (hssA) and alkylsuccinate synthase (assA), which can activate PAHs or alkanes using the fumarate addition mechanism [25–27].

The coexistence of high methane (up to 15 mM) and pore-water sulfate (up to 28 mM) concentrations in the well-ventilated Guaymas sediments create favorable conditions for the sulfate-dependent, anaerobic oxidation of methane (AOM; Additional file 1: Table S1) [12, 13]. We recovered a Methanomicrobia MAG (bin 4572_4) belonging to the archaean ANME-1 lineage, which are known to be involved in syntrophic anaerobic methane oxidation (ANME; Additional file 1: Figure S2) [28]. As previously described for other ANME members, this MAG encodes a complete pathway for methane oxidation via reverse methanogenesis (Fig. 5a) [29]. The key gene of this pathway is the mcrA gene, which encodes for the methyl–coenzyme M reductase that cleaves methane to form methyl coenzyme M [30]. This organism’s predicted McrA is closely related to previously described ANME-1 proteins (Fig. 5b). We did not detect any genes involved in electron transfer to an external electron acceptor within the Methanomicrobia MAGs; therefore, these archaea likely require a syntrophic interaction with bacterial sulfate reducers. These sulfate reducers characteristically belong to the Desulfosarcina/Desulfococcus (DSS) or Desulfobulbus-related (DSB) lineage of the Deltaproteobacteria [31, 32]. We successfully reconstructed two sulfate-reducing MAGs assigned to the DSS lineage.
that could be potential syntrophic partners of ANME-1 (Fig. 2, Additional file 1: Figure S3, Table S3).

Aside from the mcrA gene identified in ANME-1, we detected further genes in the Bathyarchaeota (bin ex4484_135) and the first representative MAGs of the GoM-Arc1 group (bins ex4572_44 and ex4484_138; Fig. 5, Additional file 1: Figure S2, Table S4) [28]. Phylogenetic analyses of these proteins revealed they are considerably divergent from those of known methane oxidizers and methanogens (Fig. 5b). In addition to having a methyl-coenzyme M reductase (subunits ABG), these archaea also contain genes encoding for enzymes required in other steps of anaerobic butane oxidation including heterodisulfide reductases (HdrABC), tetrahydromethanopterin S-methyltransferase (MtrABCDEFGH), methylenetetrahydromethanopterin reductase (Mer), N10-methenyl-H4MPT cyclohydrolase (Mch), formylmethanofuran dehydrogenases (FmdABCD), and a complete Wood-Ljungdahl pathway (Fig. 5a). However, neither of the GoM-Arc1 genomes contains genes encoding for the butyryl-CoA oxidation or beta-oxidation pathway, and the mcrA genes belonging to GoM-Arc1 are phylogenetically distinct from those previously shown to be involved in butane oxidation in Candidatus Syntrophoarchaeum spp. [33]. It has been shown that the anaerobic oxidation of hydrocarbons can be achieved by a syntrophic interaction with sulfate-reducing bacteria [33]. We obtained 15 Deltaproteobacteria MAGs belonging to a variety of taxa. The MAGs belonging to the family Desulfobacterales (bins ex472_123 and ex4572_130) contain genes that encode a type IV pilus (PilA) and extracellular cytochromes, which have been implicated in transferring electrons between archaea and bacteria [34, 35].

Community interactions
To investigate other potential biogeochemical interdependencies in the GB sedimentary communities, we mapped the ecological roles of all the microbes that were obtained (Fig. 6). This revealed several potential substrate-dependent interactions among fermentative community members and sulfur- and nitrogen-cycling organisms. Sulfate reduction pathways are encoded in Deltaproteobacteria and Archaeoglobales archaeal MAGs (Figs. 1 and 2, Additional file 1: Table S4). Furthermore, the Desulfurococcales within the archaea are predicted to be capable of S0 reduction and Bathyarchaeota MAGs encode the alpha and beta subunits of the anaerobic sulfite reductase (asrAB). Hydrothermal fluids also provide
millimolar concentrations of sulfide in GB sediments [12, 13, 19]. This sulfide could be oxidized by the dominant bacterium affiliated with the *Beggiatoaceae* (bin ex4572_84) [36], which contains genes encoding sulfide quinone reductases (*sqr*), sulfur oxidases (*soxBY*), and thiosulfate reductase (*phsA*). The *Epsilonproteobacteria* are a second phylogenetic lineage likely involved in sulfur oxidation. Based on the presence of sox genes (subunits *soxBCY*) in the reconstructed epsilonproteobacterial genomes (bins ex4484_166, ex4484_230, ex4484_65 and ex4484_4), we propose they are primarily involved in intermediate sulfur (thiosulfate) cycling.

The *Beggiatoaceae* and *Epsilonproteobacteria* appear to be capable of chemoautotrophic coupling of nitrate reduction to sulfur oxidation, consistent with previous genomic and physiological analyses [19, 36, 37]. Nitrate reduction extends beyond the *Proteobacteria*; many other bacteria belonging to *Bacteroidetes*, KSB1 and WWE1 encode nitrate reductases (*napA*) and likely require organic electron donors. In addition to utilizing hydrogen as electron donor, these organisms are predicted to take up fermentation byproducts and presumably oxidize them by denitrification; ethanol can potentially be used by *Chloroflexi* and KSB1, and acetate by *Bacteroidetes* and WWE1. Hydrogenases were often detected in nitrate reducers (Figs. 1 and 2).

We found genes encoding for Ni,Fe-Hydrogenases in *Epsilonproteobacteria*, suggesting they are able to oxidize sulfur and H₂, likely coupled to the reduction of nitrate. Furthermore, *Deltaproteobacteria* are predicted to couple sulfate reduction with the oxidation of carbohydrates, fatty acids, or petroleum-like compounds. Overall, these findings highlight the multitude of substrate-level dependencies driving nutrient cycles in GB sediments.

**Discussion**

We assembled MAGs of 115 GB-inhabiting microbes, comprising 9archael and 18 bacterial phyla, many of which are abundant in GB sediments based on diversity surveys from the same sediments [13]. The most dominant archaeal and bacterial community members belonged to a member of the archaeal ANME-1 group and the *Beggiatoaceae*, respectively (Additional file 1: Figures S8, S9). The high abundance of both taxa is expected, as both sediment locations are covered by dense *Beggiatoaceae* mats, and *Methanosarcinales* have been previously defined as abundant community members [13, 19, 36].

The reconstruction of MAGs sheds light on the metabolic potential of individual community members. Genes for the degradation of organic matter and the ability to ferment appeared to be nearly universal across GB archaea and bacteria (Figs. 1 and 2). Overall, GB microbes contained multiple genes to degrade and assimilate a broad range of substrates, including carbohydrates, peptides, petroleum compounds, and fatty acids, indicating that the utilization of varied types of sedimentary organic carbon – fresh photosynthetic biomass as well as fossil hydrocarbons - is a central function of microbial communities in these sediments. This broad access to...
different carbon sources appears highly characteristic for sediment environments, including aquifers and estuaries, suggesting that the microbial utilization of organic carbon is a central driver of nutrient cycles in marine sediments [22, 23]. Although Guaymas Basin has unique hydrothermally active sediments, these sediments share many characteristics with other sedimentary environments, such as the White Oak River estuary [38, 39]. Both sediment environments are rich in photosynthetically derived organic matter, sulfide and methane, which potentially explain similarities among these different habitats.

A unique feature of GB sediments is the high abundance of short-chain (petroleum-like) \( C_2-C_{10} \) hydrocarbons that are generated during hydrothermal pyrolysis [5]. Interestingly, we identified several potential enzymes for anaerobic hydrocarbon degradation, including putative activating enzymes involved in PAH and alkane degradation in Bacteroidetes, Chloroflexi, Deltaproteobacteria, and candidate phyla Latescibacteria and KSB1 (Fig. 2). Deltaproteobacteria are often enriched in oil-contaminated sediments and can couple the oxidation of hydrocarbons with the reduction of sulfate [40, 41]. However, to our knowledge, this is the first documentation of genes predicted to be involved in anaerobic hydrocarbon degradation in the candidate phyla Latescibacteria and KSB1. The widespread distribution of genes for hydrocarbon degradation beyond members of the Proteobacteria extends the potential metabolic diversity found in deep-sea sediments and calls for hydrocarbon degradation studies of cultivable Chloroflexi and Bacteroidetes in Guaymas Basin.

Across the archaeal genomes, ANME-1, Batharchaeota and Gom-Arc1 members are able to oxidize methane and other short-chain alkanes using McrA (Fig. 5). The ANME-1 McrA protein is likely involved in AOM that is coupled with sulfate reduction in Deltaproteobacteria [28]. For example, it has been shown that ANME-1 archaea and members of the Desulfosarcina (family Desulfo bacteraceae) are closely associated in sediments [42]. We reconstructed two MAGs that are related to the Desulfo bacteraceae and are able to reduce sulfate and encode for potential electron carriers (Fig. 2, Additional file 1: Figure S3). Therefore, we hypothesize that the ANME-1 are coupling AOM with sulfur reduction carried out in a syntrophic interaction with these Desulfo bacteraceae. This attribution has to be validated since members of this family also thrive as free-living sulfate reducers in marine sediments. At least in the case of ANME-2 archaea, the Desulfosarcina syntrophs are members of a specific 16S rRNA gene-defined clade, the SRB1a cluster [31]; since we could not retrieve 16S rRNA genes of these MAGs, further experiments are required to confirm this taxonomic affiliation. The Batharchaeota type mrcA is related to those recently described in genomes of this group from a deep aquifer that have been shown to be able to oxidize butane instead of methane [43]. The two GoM-arc1 archaeal MAGs contained novel alkyl-coenzyme M reductase (mcr) genes that clustered apart from both the ANME-1 and bathyarchaeotal mcrA genes. Based on the phylogeny of the mcrA gene and the absence of the beta-oxidation pathway [33], we conclude that the GoM-Arc1 archaea are not capable of oxidizing butane, but rather another type of short-chain hydrocarbon, using a novel pathway to feed the McrA-activated hydrocarbon into the reverse methanogenesis pathway. Overall, the phylogenetic comparison of these McrA proteins suggests that archaea use them for a broader repertoire of hydrocarbon substrates than has been realized.

Elucidating the metabolisms of individual community members allowed us to map potential biogeochemical interdependencies among the members of this hydrothermal sediment community. The fermentative organisms might metabolize complex carbon sources and thereby provide substrates to fuel the anaerobic respiration of both nitrate and sulfate. The prevalence of sulfate-dependent pathways is consistent with the high pore-water sulfate concentrations in GB sediments; from several millimolar up to full seawater strength (28 mM) [12, 13]. Notably, we found likely syntrophic relationships among both fermenting and respiring organisms, aside from the well-described interaction of ANME archaea with Deltaproteobacteria [31]. For example, the sulfur cycle was fragmented across several community members, suggesting that biogeochemical nutrient cycles are partitioned among individual community members [23]. Future experiments, including gene expression studies or amino acid tagging techniques, are needed to confirm the activity and interconnectivity of Guaymas Basin sediment-inhabiting microbes. Additionally, selecting a wider dataset, including sites spanning a broader depth and temperature profile, will help determine whether the functional diversity in this study reflects the diversity present in hydrothermal vents as a whole.

**Conclusions**

GB hydrothermal vent sediments are hotspots for microbial carbon cycling and contain high concentrations of methane and hydrocarbons, including alkanes and PAHs [5, 12, 13]. The metabolic reconstruction of 115 new microbial MAGs revealed the substrate-dependent connectivity among deep-sea inhabiting microorganisms. Mapping of the inferred ecological roles of all these organisms indicated potential biogeochemical interdependencies in organic matter utilization, hydrocarbon degradation, and respiratory sulfur and nitrogen cycling. Of particulate interest is the identification of potentially novel enzymes for hydrocarbon degradation in Chloroflexi, Bacteroidetes, and candidate phyla Latescibacteria and KSB1. Additionally, the first genomes of the GoM-arc1 archaea contained novel alkyl-coenzyme M
(mcr) genes and pathways for the oxidation of an unknown short-chain alkane. These findings extend the spectrum of hydrocarbon-degrading physiologies among deep-sea inhabiting microorganisms and call for hydrocarbon degradation studies among cultivable microbes in Guaymas Basin.

Methods
Sampling
Guaymas Basin sediment samples were collected from the Gulf of California (27° N 0.388, 111° W 24.560) at a depth of approximately 2000 m below the water surface. Sediment cores were sampled during two Alvin dives (dive 4484 core #1, December 6, 2008, and dive 4572 core #18, December 3, 2009) from a hydrothermal mat area at the base of Mat Mound and from a 200-m distant hydrothermal area, termed Marker 27 (Additional file 1: Table S1). Dense mats of *Gammmaproteobacteria* of the family *Beggioatoaceae* covered both sites, with a white mat dominating at site 4484 and an orange mat at site 4572. Intact sediments from both dives were collected using polycarbonate cores (45 cm in length, 6.25-cm interior diameter), subsampled into centimeter layers under N₂ gas in the ship’s laboratory and immediately frozen at −80 °C. Sediment subsamples for DNA isolation were taken from depths 0–1 and 3–4 cm in core 4484-1 and 0–3 and 12–15 cm in core 4572-18. Metadata for dive 4484 and 4572, including details on the geochemistry (i.e., methane concentrations, dissolved organic carbon concentrations, sulfate and sulfide concentrations) as well as thermal profiles of the sampling site, are available to compare microbial community composition across sediment cores [12, 19, 21]. For 4484-1, full metadata are included in Dowell et al. 2016. For 4572-18, full metadata are included in McKay et al. 2016. Pictures of the sampling locations are included in Teske et al. 2016.

Metagenomic sequencing
Total DNA from 10 g of sediment from each of the four samples (see above) was extracted using the MoBio PowerMax soil kit. DNA concentrations were measured using a Qubit™ 3.0 Fluorometer, and a final concentration of 10 ng/µl of each sample (using a total amount of 100 ng) was used to prepare libraries for paired-end Illumina (HiSeq 2500) sequencing. Illumina library preparation and sequencing was performed by the GSAF (Genome Sequencing and Analysis Facility) at the University of Texas at Austin. Sequencing was performed on an Illumina HiSeq 2500 with the following specifications: high throughput run mode, run type paired end 2 × 125 bp, 6 × 4.0E8 target reads (millions), insert size approximately 360–420 bp and ~5% PhiX control spike-in. This sequencing approach provided a total of ~242 gigabases of sequencing data (411,732,022/520,750,012/552,634,962/452,269,130 reads from the sediment samples 4484 0–1 cm/4484 3–4 cm/4572 0–3 cm/4572 12–15 cm, respectively).

Raw Illumina shotgun genomic reads were separated from Illumina artifacts by removing the adaptors and DNA spike-ins from the forward and reverse reads. Reads with an average quality score < Q20 and a read length < 50 bps were removed using cutadapt [44]. Afterwards, reads were interleaved using interleave_fasta.py and the interleaved sequences were trimmed using Sickle with default settings [45]. The script for interleave_fasta.py can be found at https://github.com/jorvis/biocode/blob/master/fasta/interleave_fasta.py. Metagenomic reads from dive 4484 and 4572 (concatenated per depth profile for better coverage) were individually assembled using IDBA-UD using the following parameters: -pre_correction, -mink 75, -maxk 105, -step 10, -seed_kmer 55 [46]. This yielded a total of 3,139,208 and 159,5687 scaffolds from sample 4484 and 4572, respectively, including scaffolds with a minimum and maximum scaffold length ranging from 200 to 177,401 bp (maximum scaffold length for sample 4484 and 4572: 177,401 and 133,414, respectively).

Metagenomic binning was performed on assembled samples from dive 4484 and 4572 by calculating tetranucleotide frequencies of scaffolds with a minimum length of 5000 bp (including a total of 58,488 scaffolds with a total length of 591,330,171 bp) [47]. The resulting Self-Organizing Maps (ESOM) were manually sorted and curated (Additional file 1: Figure S1) [47]. Metagenomic binning was enhanced by incorporating reference genomes as genetic signatures for the assembled contigs into ESOM [47, 48]. Thereby, we assembled 77 bacterial and 38 archaeal metagenomic assembled genomes (MAGs) with a completeness above 50%. After binning, MAGs were linked to the original sediment samples based on their unique scaffold ID. CheckM was employed to evaluate the accuracy of the binning approach by determining the percentage of completeness and contamination (Additional file 1: Table S3) [49]. Contaminants that were identified based on their phylogenetic placement (wrong taxonomic assignment compared to the average taxonomic assignment of the genes assigned to each bin), GC content (>25% difference compared to the mean of all scaffolds assigned to each bin), or confidence level (>25% differences compared to the mean of all scaffolds assigned to each bin) were manually removed from each MAGs.

Genome coverage
To determine the relative abundance of each MAG across the four sequenced Guaymas Basin sediment samples, we mapped scaffolds from all MAGs against the original metagenomic sequencing reads using BWA using default
settings [50]. To detect exact matches to the original sequencing data, we only considered matches where the complete sequence of the raw sequence read (125 bp) matched the MAG. The relative abundance of each MAG was calculated by normalizing the recruited reads per MAG by the genome size, accounting for differences in sampling depth of the respective metagenome and then multiplying by 1,000,000. A total of 31,363,251 archaeal and 116,888,222 bacterial reads mapped back to the original metagenomes and represented ~15% (archaea) and ~49% (bacteria) of the sequenced community.

**Gene calling, taxonomic assignment, and functional characterization**

Gene calling and taxonomic assignment for the four metagenomic samples and individual MAGs was performed using the Joint Genome Institute-Production Genomics Facility (IGI-PGF) integrated microbial genomes with microbiome (IMG/M) system. The IMG output was linked to the MAGs by their unique scaffold ID, which was also used to extract protein sequence information for further analyses. Additionally, gene calling for individual MAGs was performed using RAST (Rapid Annotation using Subsystem Technology) [51, 52]. For RAST, individual MAGs were uploaded using the Network-Based SEED API using the command svr_submit_RAST_job, selecting the RAST gene caller method.

For a further functional characterization, individual MAGs were analyzed using the IMG/M systems output, the SEED subsystems, KAAS (KEGG Automatic Annotation Server), and the Carbohydrate-Active enZYMes (CAZy) database [53, 54]. For the KAAS- and CAZy-based analysis, concatenated protein fasta sequences of each MAG were uploaded to the KAAS and dbCAN webservers using the metagenome setting for KAAS (parameters: GHOSTX, genes dataset, SBH assignment method) and default settings for dbCAN.

Additionally, we searched for key metabolic genes using custom blast and hmmer databases using previously defined thresholds [23]. Therefore, we manually curated a reference blastp database including metabolic genes by searching the KEGG and NCBI databases for pathways and corresponding genes of interest. This reference database was screened against the concatenated protein fasta sequences from the MAGs using blastp (e-value threshold of 1e−20) [55]. Additionally, we utilized a published hmmer database using hmmsearch and custom bit score thresholds [23]. Hydrogenases were extracted from the genomes using hmmsearch (e-value cut-off of 1e−20), and hits were confirmed using a web-based search using the hydrogenase classifier HydDB [56]. Other positive hits of the blast or hmmer search were manually confirmed using a NCBI-based protein blast search [55].

The MAGs on average contained 1285 (archaea) to 2043 (bacteria) protein-coding genes, and of these, ~73% could be functionally assigned (Additional file 1: Table S3).

The core marker genes for processes shown in Figs. 1 and 2 as well as Additional file 1: Figures S3, S4, and S5 are listed in Additional file 1: Table S4. This table includes the information on the used databases as well as thresholds used for the functional annotation. Thresholds for the hmmer search were used as previously described [23]. Key enzymes and subunits were identified using hmmer searches, protein blast databases, and the KAAS and IMG/M systems. General pathway searches using the KAAS and IMG/M systems were used to confirm the presence of corresponding metabolic pathways in each of the bins. Functional processes were only considered if key enzymes/subunits were identified in multiple databases and when >50% of the corresponding pathway components were detected in a genome.

**Phylogenetic analyses**

Phylosift was used to extract marker genes for the phylogenetic placement of the assembled metagenomic bins [57]. These marker genes consist of up to 15 syntenic ribosomal protein genes that have been demonstrated to undergo limited lateral gene transfer [rpl2, 3, 4, 5, 6, 14, 15, 18, 22, 24 and rpS3, 8, 10, 17, 19] [58]. This gene set was derived from a reference database as detailed in [59]. To confirm the placement of the Candidatus Stahlbacteria as a separate phylum, we extracted up to 37 marker genes included in Phylosift for a more robust phylogenetic placement. These single-copy protein-coding markers include rpS2, rpS3, rpS5, rpS7, rpS8, rpS9, rpS10, rpS11, rpS12, rpS13, rpS15P, rpS17, rps19, rpl1, rpl2, rpl3, rpl4, rpl5, rpl6, rpl11, rpl13, rpl14b, rpl15, rpl16, rpl18P, rpl22, rpl24 rpl25, rpl29, IF-2, phenylalanyl-tRNA synthetase alpha and beta subunit, tRNA pseudouridine synthase B, porphobilinogen deaminase and the ribonuclease HII. To search for ribosomal protein sequences, all MAGs (fasta files) were used as an input in Phylosift, which was used with default parameters. Moreover, we included sequences from bacterial reference strains for phylogenetic analyses [60]. Amino acid alignments of the individual ribosomal protein genes were generated using MAFFT and manually curated [61]. For the extended phylogeny using 37 marker genes, we used the concatenated protein alignment provided by Phylosift. Afterwards, the curated alignments of the ribosomal proteins were concatenated for further phylogenetic analyses. For the ribosomal protein alignments,
Additional file

Additional file 1: Supplementary Tables S1-S5 and Figures S1-S10. Tables S3 and S4 are provided separately. Table S3. Summary statistics of assembled archaeal and bacterial MAGs. Metagenome-assembled genomes (MAGs) assembled from dive 4484 and 4572. Providing information about the taxonomic affiliation (ribosomal protein phylogeny), completeness (CheckM) and genome statistics. The completeness lists the CheckM results, including the number of evaluated markers, the numbers of times a marker was found within a respective genome (numbers ranging from 0 to 5+) and the degrees of completeness, contamination, and heterogeneity. Summary statistics include the GC content (%), genome length (Mb), average coverage (based on read coverage of the individual contigs), number of called genes and average gene length (bp). Table S4. Functional analysis of assembled archaeal and bacterial MAGs. Functional gene analysis of individual archaeal and bacterial MAGs recovered from dive 4484 and 4572 based on HMMER, KAAS, KEGG, and blastp analyses of marker genes for specific pathways involved in carbon (C), nitrogen (N), and sulfur (S) cycling as well as several additional pathways. Presence/absence of genes are listed as: Presence: >1 (red), Absence: 0 (no color). (ZIP 2404 kb)

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Availability of data and materials

All sequence data and sample information are available at NCBI under BioProject ID PRJNA362212. Accession numbers for individual MAGs are provided in Additional file 1: Table S3.

Authors’ contributions

BJB and AT conceived and designed the study. KWS and ND processed the data and reconstructed the genomes. ND, KWS, and BJB analyzed the functional annotations and the metabolisms. All authors were involved in writing the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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