Bacteria and Archaea Synergistically Convert Glycine Betaine to Biogenic Methane in the Formosa Cold Seep of the South China Sea

Lingyan Li,a,b Wenting Zhang,a,b Shengjie Zhang,a,b Lei Song,c Qinglei Sun,d Huan Zhang,d Hua Xiang,a,b Xiuzhu Donga,b

aState Key Laboratory of Microbial Resources, Institute of Microbiology, Chinese Academy of Sciences, Beijing, China
bUniversity of Chinese Academy of Sciences, Beijing, China
cChina General Microorganism Culture Collection Center, Institute of Microbiology, Chinese Academy of Sciences, Beijing, China
dInstitute of Oceanology, Chinese Academy of Sciences, Qingdao, China

Lingyan Li, Wenting Zhang, and Shengjie Zhang contributed equally to this work. Author order was determined in order of the time participating in the project.

ABSTRACT Cold seeps are globally widespread seafloor ecosystems that feature abundant methane production and flourishing chemotrophic benthic communities. Chemical evidence indicates that cold seep methane is largely biogenic; however, the primary methane-producing organisms and associated pathways involved in methanogenesis remain elusive. This work detected methane production when glycine betaine (GBT) or trimethylamine (TMA) was added to the sediment microcosms of the Formosa cold seep, South China Sea. The methane production was suppressed by antibiotic inhibition of bacteria, while GBT was accumulated. This suggests that the widely used osmoprotectant GBT could be converted to cold seep biogenic methane via the synergistic activity of bacteria and methanogenic archaea because archaea are not sensitive to antibiotics and no bacteria are known to produce ample methane (mM). 16S rRNA gene diversity analyses revealed that the predominant bacterial and archaeal genera in the GBT-amended methanogenic microcosms included Oceanirhabdus and Methanococcoides. Moreover, metagenomic analyses detected the presence of grdH and mtgB genes that are involved in GBT reduction and demethylation, respectively. Two novel species were obtained, including bacterium Oceanirhabdus seepicola, which reduces GBT to TMA, and a methanogenic archaeon, Methanococcoides seepicolus, which produces methane from TMA and GBT. The two strains reconstituted coculture efficiently converted GBT to methane at 18°C; however, at 4°C addition of dimethylglycine (DMG), the GBT demethylation product, was necessary. Therefore, this work demonstrated that GBT is the precursor not only of the biogenic methane but also of the cryoprotectant DMG to the microorganisms at the Formosa cold seep.

IMPORTANCE Numerous cold seeps have been found in global continental margins where methane is enriched in pore waters that are forced upward from sediments. Therefore, high concerns have been focused on the methane-producing organisms and the metabolic pathways in these environments because methane is a potent greenhouse gas. In this study, GBT was identified as the main precursor for methane in the Formosa cold seep of the South China Sea. Further, synergism of bacteria and methanogenic archaea was identified in GBT conversion to methane via the GBT reduction pathway, while methanogen-mediated GBT demethylation to methane was also observed. In addition, GBT-demethylated product dimethyl glycine acted as a cryoprotectant that promoted the cold seep microorganisms at cold temperatures. GBT is an osmoprotectant that is widely used by marine organisms, and therefore, the GBT-derived methanogenic pathway reported here could be widely distributed among global cold seep environments.

Citation Li L, Zhang W, Zhang S, Song L, Sun Q, Zhang H, Xiang H, Dong X. 2021. Bacteria and archaea synergistically convert glycine betaine to biogenic methane in the Formosa cold seep of the South China Sea. mSystems 6:e00703-21. https://doi.org/10.1128/mSystems.00703-21.

Editor Zarath M. Summers, ExxonMobil Research and Engineering

Copyright © 2021 Li et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license.

Address correspondence to Lingyan Li, lilingyan@im.ac.cn, or Xiuzhu Dong, dongsx@im.ac.cn.

Received 7 June 2021
Accepted 17 August 2021
Published 7 September 2021
Cold seeps are localized areas within cold marine systems that exhibit considerable methane release to seafloor fluids and are therefore termed methane seeps. Methane seeps are enriched in methane-metabolizing chemotrophic bacteria and archaea that can transform the carbon and chemical energy stored in methane to biomass and biological energy, respectively, in turn supporting flourishing faunal assemblages (1–4). In addition, cold seeps represent large methane sources that can contribute to global warming. Consequently, the discovery and investigation of cold seeps 40 years ago have led to a focus on the methane-metabolizing organisms within these systems, including the anaerobic methanotrophogenic archaea (5–8).

The stable carbon isotope signatures (δ¹³C) of the cold seep methane suggest biogenic sourcing in some regions, as observed for the methane released from the Sonora cold seep in the Guaymas Basin that exhibits δ¹³C values of about −63%, indicating a presumably dominant proportion of biogenic mixed with abiotic methane (e.g., thermogenic methane) (9–11). A cold seep at Formosa Ridge (also termed site F) is located on the northeastern corner of the continental slope of the South China Sea (SCS) (12), where rich communities of benthic invertebrates are found, indicating that this is an active cold seep. The stable carbon isotope signatures (δ¹³C) of Formosa cold seep Bathymodiolus platifrons tissues containing methanotrophic symbionts are −70.3% on average (13). Moreover, methane of the Haima cold seep in SCS also exhibits δ¹³C values of −72 to −99%. These observations also implicate the presence of biogenic methane in the SCS cold seeps.

Methylotrophic methanogenesis and, specifically, methane production from trimethylamine (TMA) has been conventionally viewed as the primary pathway of ocean ecosystem methanogenesis. Importantly, sulfate-reducing bacteria (SRB) and other bacteria, except for some acetogens (14, 15), in anoxic marine sediments are incapable of metabolizing TMA (16), leaving TMA metabolism solely to methanogens without competition. Quaternary amines, glycine betaine (GBT), and choline are ubiquitous osmolytes used by marine microorganisms and benthic invertebrates and are all precursors of TMA. These compounds are assumed to contribute as metabolites to as much as 90% of total methane production in intertidal coastal sediments (17). In addition, other methyl compounds like methylphosphonate could be used by Vibrio species to produce methane. Methane production was also found in oxic seawater of the western North Pacific and its marginal seas (18). TMA-utilizing methanogens in marine ecosystems primarily belong to the genera Methanococcales and Methanosarcina (19, 20). However, only a few methanogenic species have been reported to directly use quaternary amines to produce methane, including Methanococcales sp. strain Q3C and Methanolobus sp. strain B1d (21). In addition, other Methanococcales strains isolated from marine sediments have been shown to produce methane from choline, dimethyl ethanolamine (DMEA), and GBT (22, 23). A nonpyrrolysine trimethylamine/corrinoid methyltransferase (MttB) homolog in Desulfotobacterium hafniense (that is capable of growing on GBT) has been shown to convert GBT and cobalamin to dimethylglycine (DMG) and methylcobalamin and was thus defined as a GBT methyltransferase (MtgB) (24). Similarly, an MtgB ortholog from Methanolobus sp. B1d was determined to transfer the GBT methyl group to coenzyme M, thereby playing a key role in GBT-dependent methylotrophic methanogenesis (25).

Most methanogens are incapable of efficiently converting GBT and choline to methane but first require the conversion of GBT or choline to TMA by bacteria. Coastal marine organisms typically accumulate GBT, with intracellular GBT being observed at concentrations of up to 1 M in some microorganisms living in hypersaline environments (26). Microcosm studies using coastal marine sediments have estimated that up to 90% of methane emitted from marine ecosystems can be produced from GBT and other
structurally related quaternary amine compounds (17, 27). In addition, TMA N-oxide (TMAO) is an abundant compound in marine eukaryotic cells (28, 29) that is another potential precursor of TMA. Together, TMAO, GBT, and choline all can be converted to TMA by TMAO and dimethyl sulfoxide (DMSO) reductase (TorA and DMSOR), GBT reductase (GrdH), or choline-TMA lyase (CutCD), respectively.

Molecular studies, including 16S rRNA gene diversity surveys and carbon-isotope coupled metagenome-assembled genome analysis, have shown that *Pelobacter* and *Methanococcoides* can syntrophically convert choline to methane in coastal saltmarsh sediments (30), while members of the novel family *Candidatus* “Betainaceae” fam. nov. that are affiliated with *Clostridiales* can convert GBT to TMA (26). In addition, the sulfur-reducing bacterium *Desulfuromonas acetoxidans* can degrade GBT to produce TMA and acetate via GBT reductase; it also converts GBT to methane in synergy with a methylo trophic methanogen (17). Nevertheless, the methanogenic archaea that produce methane and the bacterial taxa that convert GBT/choline to the methanogenic precursor TMA in cold seeps remain largely unknown.

In this study, GBT was used as the sole carbon source to obtain the Formosa cold seep microcosms, which efficiently converted GBT to methane. Species diversity and metagenomic analyses revealed the predominant methanogenic archaea and GBT-degrading bacterial species and the genes encoding GBT reductase (GrdH) and methyltransferase (MtgB) that are involved in the two primary anaerobic GBT degradation pathways, GBT reduction and demethylation, respectively. The isolated bacterial strain (ZWT) and methanogenic strain (LLY) both represent new species, respectively, and converted GBT to TMA and methane, and a reconstructed coculture of the two strains was able to convert GBT to methane at lower temperatures corresponding to in situ conditions. Remarkably, dimethylglycine (DMG), a GBT demethylation product, promoted growth of the two strains at lower temperatures, either in mono- or cocultures. Overall, this work determined that the widely used osmolyte GBT is the precursor for methane production in the Formosa cold seep and also of cryoprotectant for cold seep microorganisms.

**RESULTS**

**Glycine betaine supports biogenic methane production in the Formosa cold seep.** To explore the microorganisms and metabolic pathways responsible for the biogenic methanogenesis in cold seeps, the reductive cold seep sediments at 1.165 m depth of the active site of Formosa Ridge (Fig. 1a; 22°06’89.05”N and 119°17’916.384”E) were sampled using a gravity core sampler during the Kexue 2019 expedition.

Cold seep sediment samples were used to establish methane production microcosms via inoculation into artificial seawater containing trace casein and amended either with methanogenic substrates or without substrates. After incubation at 18°C under N₂ and CO₂ headspaces (unless indicated otherwise), methane was produced from all of the sediment microcosms after 9 weeks of incubation. Six- and 4-fold higher methane contents were produced from the microcosms containing TMA and GBT than those from artificial seawater, but similar lower levels of methane were observed in those amended with choline, acetate, and H₂ plus CO₂ (Fig. 1b). This suggests that GBT and its reductive product, TMA, could be the major precursors for biogenic methanogenesis in the cold seep. In addition, more methane was produced from the upper sediment layer (0 to 20 cm below seafloor [cmbsf]) than the deeper layer (20 to 60 cmbsf), indicating that biogenic methane is mainly produced in the upper seafloor layer. To discern whether GBT-supported methanogenesis was mediated by methanogenic archaea alone or via a syntrophy of methanogenic archaea and bacteria in cold seep sediments, ampicillin and kanamycin were used to inhibit bacterial activities. Antibiotic amendment almost completely inhibited methane emissions, wherein only 0.05 mmol CH₄/liter and 1.65 mmol/liter GBT were observed in the microcosm without addition of methanogenic precursors. Therefore, 97% of the methane production from GBT was estimated based on the 2.1 mmol CH₄/liter emitted from the noninhibited microcosms. Antibiotic inhibition of methane production indicates that bacteria that
presumably convert GBT to TMA synergistically with methanogenic archaea contribute to biogenic methanogenesis in cold seeps.

In addition, carbon isotope fractionation assays indicated that the sediment-released CH₄ possessed a similar but slightly heavier carbon isotope signature (δ¹³C, 241.82 to 261.49%, depending on CH₄ concentrations) than that from the GBT-amended microcosms (247% to 271.56%) (Fig. 1b, inset). These values were much lighter than the δ¹³C of acetate-derived CH₄ (29% to 35%) and between CO₂ reduction-produced methane (28% to 79%). The carbon isotope fractionation values of the cold seep methane further suggested that biogenic methane in the system is most likely derived from GBT or its reductive product, TMA.

Uncultured Clostridiales spp., Methanococcoides spp., and the genes involving GBT metabolism are enriched in GBT-amended cold seep microcosms. GBT was used as the sole carbon source to enrich the cold seep microorganisms that convert GBT to CH₄. The upper layers from the seep sediments were incubated in 20 mM GBT for 81 days at 18°C, followed by measurement of CH₄ production to identify GBT-dependent methanogenic microcosms. 16S rRNA gene diversity analyses revealed that the dominant bacterial genera in the GBT microcosms included Vallitalea, a Gram-negative halophile, Tyzzerella and Oceanirhabdus within the order Clostridiales (family Clostridiaceae), and the facultatively anaerobic Draconibacterium. In contrast, Pseudomonas, Sulfurovum, and unidentified Clostridiales were abundant in in situ sediments (Fig. 2a). Archaeal communities also markedly shifted after GBT enrichment, with dominant organisms becoming methanogens like Methanococcoides (<6% in sediment to 22.50% in microcosms) and Methanobacterium (<0.1% in sediment to 18.83% in microcosms) in addition to uncultured Bathyrhoea (<0.1% in sediment to 30.49% in microcosms). In contrast, ANME-2a (28.91% in sediment) and ANME-1b (31.25% in sediment) groups and Thaumarchaeota group B (24.22% in sediments) were prevalent in in situ sediments but disappeared in the GBT enrichment after 7 weeks (Fig. 2b). After 81 days of incubation, the uncultured Bathyrhoea also disappeared, and Methanococcoides increased to >90% relative abundance. The archaeal community shifts were consistent with methane production in the microcosms.

Metagenomic analyses were then conducted on cold seep sediments and GBT microcosms using Illumina HiSeq-based shotgun metagenomic sequencing. The assembled contigs were used for functional annotations, and proteins pertinent with
GBT reduction (GrdH and GrdB) and its transporter (OpuD) were further analyzed for phylogenetic clustering with the respective functionally defined proteins (Fig. S1 in the supplemental material). In both the in situ sediments and GBT microcosms, genes or gene clusters involved in the two major GBT anaerobic degradation pathways, GBT reduction (Fig. 3a) and GBT demethylation (Fig. 3b), were enriched. Genes involved in the GBT reduction pathway included *grdHI*, which encodes GBT reductase complex B that specifically reduces GBT to TMA, and acetyl phosphate, *opuD*, which encodes a transporter for GBT, and the structurally analogous *grdABCD* encoding a reductase complex that reduces glycine, sarcosine, or GBT. In addition, genes encoding thioredoxin and its reductase that could provide reducing equivalents to GBT reduction were also detected. Overall, 26 contigs possessing *grdHI* were obtained, and these genes were organized into 6 cluster types (Fig. 3a). Of these contigs, some (like those in types I, III, and V) were enriched by about 100-fold in GBT microcosms. In addition, 503 contigs carrying the *mtgB* gene cluster, which is likely involved in GBT demethylation, were identified. It is worth noting that *mtgB* (encoding the GBT corrinoid protein-CoM methyltransferase) and *mtgC* (encoding the GBT corrinoid protein) were clustered with more versatile GBT transporters than those clustered with *grdHI* genes. In addition to *opuD* clustering with *grdHI*, *opuABC* (encoding an osmoprotectant transport system), and another two potential GBT transporters (*proWX* encoding GBT/proline transporters and *ABC.AP* encoding ABC transporters) were also identified. The identified *mtgB* genes were also organized as six cluster types, with types I, IV, V, and VI being enriched by 10- to nearly 300-fold (Fig. 3b). This suggests that two GBT metabolic processes, GBT reduction to TMA and demethylation to dimethylglycine, could co-occur in the cold seep environments. In contrast, genes encoding TMAO reductase (TorA) or choline-TMA lyase (CutCD) were not found in either the in situ sediments or GBT microcosms, thereby rendering TMA production from either TMAO or choline unlikely.

Sequence homology analysis indicated that the 13 types of identified *grdH* were associated with numerous organisms, with 38 *grdH* orthologs detected in the cold seep primarily associated with unclassified *Firmicutes*, *Proteobacteria*, *Spirochaetia*, and *Fusobacteria* (Fig. S2a). The 503 enriched orthologs were mainly associated with *Firmicutes*, *Chloroflexi*, *Proteobacteria*, and *Planctomycetes* (Fig. S2b), while those from *Euryarchaeota* were underrepresented. It is worth noting that the MtgB phylogeny was not topologically congruent with that for the species carrying the corresponding MtgB orthologs, suggesting the presence of frequent horizontal gene transfer in cold seep sediment systems. Given that only a small portion of the cold retrieved *mtgB* from methanogenic archaea, we predicted that methanogen-
mediated GBT-derived methanogenesis could have little contribution to the cold seep biogenic methane.

**Oceanirhabdus seepicola** sp. nov., a novel bacterium isolated from cold seep microcosms, reduces GBT to TMA at 18°C. The bacterial strain ZWT was isolated from GBT-enriched cold seep microcosm at day 41 using GBT as the sole carbon source. Strain ZWT converted 20 mM GBT to TMA stoichiometrically, in addition to using glycine, glucose, and maltose as sole carbon sources (Table S1). Strain ZWT exhibited the highest 16S rRNA sequence identity (94%) with *Oceanirhabdus sediminicola* NH-JN4 (Fig. 4a) within the *Clostridiaceae* family. Thus, strain ZWT was identified as a novel species of the genus *Oceanirhabdus* as *Oceanirhabdus seepicola* sp. nov. for noting a seep inhabitant and was deposited in the China General Microbiological Culture Collection Center under accession no. CGMCC 1.17897 and the Japan Collection of Microorganisms under accession no. JCM 34232.

The genome of *Oceanirhabdus seepicola* sp. nov. (JAGSOJ000000000; Table S2) possesses two gene clusters encoding GBT reductases (Fig. 4b). One cluster encodes the GBT-specific reductase complex B, including *grdI*, *grdH*, and *grdB* that encode the alpha, beta, and gamma subunits and *opuABDC* that encodes the GBT transporter. The other cluster comprises *grdE* and *grdAB* that encode glycine/sarcosine/GBT reductase complex components A and B. In addition, genes encode trimethylamine permutase (*GM000616*, *GM002195*, and *GM002197*) were found in the genome. Although the genome contains a gene encoding the choline-TMA lyase-activating enzyme, the gene encoding choline-TMA lyase (*CutC*) was not identified. Therefore, strain ZWT carries the GBT reductase genes that were detected in the cold seep metagenomes.
Consistent with the genomic analyses, *O. seepicola* ZWT can use GBT as the sole carbon source, which supported a growth rate of 0.189 per hour, but it also converts GBT to equimolecular concentrations of TMA at a degradation rate of 0.01 mmol/C1 h at 18°C (Fig. 4c). In addition, *O. seepicola* ZWT can grow with casein as the sole carbon and nitrogen sources, but not choline. Compared to *Oceanirhabdus sediminicola* NH-JN4, *O. seepicola* ZWT grew at a lower temperature range (4 to 30°C), and optimally between 18 to 22°C, and was able to use GBT and glucose (Table S1). Therefore, the distinct phenotypic characteristics of strain ZWT also supported its independent species status along with lower 16S rRNA sequence homology.

*Methanococcoides seepicolus* sp. nov. isolated from the GBT microcosms of cold seep encodes a GBT reductase gene cluster and efficiently reduces GBT to TMA at 18°C. (a) Neighbor-joining phylogenetic tree based on 16S rRNA genes of strain ZWT and its closest relatives. GenBank accession numbers are given in parentheses. Percentages of bootstrap support are shown at branch nodes. Scale bar indicates 2% sequence divergence. (b) Gene clusters involved in GBT reduction identified in the strain ZWT genome. Gene annotations are the same as in Fig. 3a, and gene numbers are shown underneath individual genes. The genome locations of the genes involved in GBT reduction are also shown. (c) Strain ZWT was grown in 20 mM GBT at 18°C, and growth was monitored by measuring OD600 values over 1-h intervals. In addition, GBT and TMA contents in cultures were measured using high-performance liquid chromatography (HPLC). Three culture replicates were assayed for each measurement and the averages and standard deviations are shown.

FIG 4

Consistent with the genomic analyses, *O. seepicola* ZWT can use GBT as the sole carbon source, which supported a growth rate of 0.189 per hour, but it also converts GBT to equimolecular concentrations of TMA at a degradation rate of 0.01 mmol-h⁻¹ at 18°C (Fig. 4c). In addition, *O. seepicola* ZWT can grow with casein as the sole carbon and nitrogen sources, but not choline. Compared to *Oceanirhabdus sediminicola* NH-JN4, *O. seepicola* ZWT grew at a lower temperature range (4 to 30°C), and optimally between 18 to 22°C, and was able to use GBT and glucose (Table S1). Therefore, the distinct phenotypic characteristics of strain ZWT also supported its independent species status along with lower 16S rRNA sequence homology.

*Methanococcoides seepicolus* sp. nov. isolated from the GBT microcosms produces methane from TMA and through GBT demethylation. A methanogenic archaeon was also isolated from the GBT microcosm developed from cold seep sediments at day 41. A methane-producing cocci-like strain LLY was obtained that used GBT as its sole carbon source. Strain LLY (CGMCC 1.17896) exhibited 99% 16S rRNA gene similarity to that of *Methanococcoides alaskenses* and *Methanococcoides burtonii* (Fig. 5a) and was thus identified as a member of the genus *Methanococcoides*. The complete genome of *Methanococcoides* LLY (JAGSOI000000000; Table S2) was then sequenced, which exhibited an average nucleotide identity (ANI) of 91% to the genome of *M. burtonii*, while a genome for *Methanococcoides alaskenses* is unavailable. Therefore, the novel species *Methanococcoides seepicolus* sp. nov. for noting a seep inhabitant was proposed for strain LLY.

The *M. seepicolus* genome contained the entire gene suite involved in methylotrophic methanogenesis, in addition to those pertinent to methanogenesis from GBT, which include mtgB and mtgC that encode GBT corrinoid protein-coenzyme M methyltransferase and GBT corrinoid protein, respectively, as well as the gene encoding the BCCT family of glycine/GBT ABC transporter permeases. The MtgBC protein complex catalyzes the transfer of the GBT methyl group to CoM, which is then reduced to methane by CoM reductase. In addition, GBT is demethylated to dimethylglycine (DMG). Genes encoding GBT or choline ABC transporters were also identified in the genome (Fig. 5b).

Unlike *M. alaskenses* and *M. burtonii*, *M. seepicolus* LLY can grow on and convert 1 mol of GBT to roughly 0.75 mol of methane (Fig. 5c), in contrast to 3 mols CH₄ produced per mol of TMA. This suggests that strain LLY only uses one of the three methyl groups of GBT to produce CH₄ via the GBT demethylation pathway, and it is consistent with the GBT methyltransferase MtgBC genes in the genome. In contrast, a lower methanogenesis...
The rate was identified when using GBT compared to that from TMA (Fig. 5c), suggesting that TMA is the preferential substrate of *M. seepicolus*. In addition to GBT and TMA, *M. seepicolus* also used methanol, but not choline, to produce methane. Cells grew over a temperature range of 0 to 25°C, and optimally at 18°C, which differed from growth characteristics of *M. alaskenses* and *M. burtonii* and confirmed the distinct species status (Table S3) along with the observed lower-genome ANI.

*Oceanirhabdus seepicola* and *Methanococcoides seepicolus* syntropically convert GBT to methane. To investigate how the two cold seep isolates synergistically convert GBT to methane, 10% of the mid-exponential cultures for each of *O. seepicola* and *M. seepicolus* were inoculated into basal medium containing 20 mM (0.45 mmol) GBT as the sole carbon source. After 15 days of incubation at 18°C, GBT was completely degraded at rates of 0.24 mmol/C1 day⁻¹ at day 1 and at 0.025 mmol/C1 day⁻¹ between days 2 and 15. TMA concentrations also peaked in the first day and then decreased to 0.05 mmol at a rate of 0.02 mmol/C1 day⁻¹. Methane production initiated at day 2 and continued until reaching 1.1 mmol at a rate of 0.097 mmol/C1 day⁻¹, and with a final CH₄ yield of about 2.8 per 1 GBT degraded (Fig. 6a). This indicates that GBT is reduced by *O. seepicola* to TMA but not by *Methanococcoides* via GBT demethylation, while the TMA is then converted to CH₄ by *M. seepicolus*. It is worth noting that cocultures exhibited lower rates of either GBT degradation or methane production than monococultures. Thus, inhibiting factors were likely present in the coculture.

We then compared expressions of the key genes involved in GBT metabolism in monococultures and in coculture. Total RNA was extracted from cocultures and GBT monococultures of *O. seepicola* and *M. seepicolus* were inoculated into basal medium containing 20 mM (0.45 mmol) GBT as the sole carbon source. After 15 days of incubation at 18°C, GBT was completely degraded at rates of 0.24 mmol·day⁻¹ at day 1 and at 0.025 mmol·day⁻¹ between days 2 and 15. TMA concentrations also peaked in the first day and then decreased to 0.05 mmol at a rate of 0.02 mmol·day⁻¹. Methane production initiated at day 2 and continued until reaching 1.1 mmol at a rate of 0.097 mmol·day⁻¹, and with a final CH₄ yield of about 2.8 per 1 GBT degraded (Fig. 6a). This indicates that GBT is reduced by *O. seepicola* to TMA but not by *Methanococcoides* via GBT demethylation, while the TMA is then converted to CH₄ by *M. seepicolus*. It is worth noting that cocultures exhibited lower rates of either GBT degradation or methane production than monococultures. Thus, inhibiting factors were likely present in the coculture.

We then compared expressions of the key genes involved in GBT metabolism in monococultures and in coculture. Total RNA was extracted from cocultures and GBT monococultures of *O. seepicola* and *M. seepicolus* at the respective mid-exponential phases, followed by estimation of transcript abundances for corresponding genes via quantitative reverse transcription-PCR (RT-PCR). It was found that the *O. seepicola* *grdH* exhibited about 60-fold higher expression levels than *M. seepicolus* *mtgB* in coculture (Fig. 6b). Similarly, *O. seepicola* *opuD* (encodes a GBT transporter) exhibited about 1,000-fold higher transcription than the archaeal ortholog. In contrast, the *O. seepicola* *grdH* and *opuD* all exhibited decreased transcriptional expression in coculture.
compared to in monococulture (Fig. 6c), suggesting that they were suppressed by unknown factors in the coculture.

*M. seepicolus* mtgB maintained similar expression levels in the mono- as in cocultures, while *mttB* (encoding trimethyltransferase) was increasingly expressed in the cocultures compared to the monococultures growing in GBT (Fig. 6d). Collectively, the gene expression profiles support that *O. seepicola* is the major player in GBT metabolism within the coculture by catabolizing GBT reduction to TMA, while *M. seepicolus* converts TMA to methane instead of demethylating GBT to dimethylglycine, most likely attributed to the lower competitiveness of the archaeon for attaining GBT when in coculture with the bacterial partner.

**The GBT demethylation product dimethyl glycine acts as a cold protectant for both the bacterium and archaeon.** To mimic in situ temperatures, *M. seepicolus* was grown at 4°C with GBT and TMA individually. Unexpectedly, though higher methane production rates were observed from TMA (0.03 mmol·day⁻¹) than from GBT (0.01 mmol·day⁻¹), a much longer lag time (25 days) was observed for the TMA culture than the GBT culture (5 days) (Fig. 7a), while the lag time difference when grown on the two substrates was not observed for 18°C cultures (Fig. 5c). Given that GBT demethylation will produce dimethylglycine (DMG), which is a cold protectant for *Bacillus* (31, 32), DMG was amended into the TMA culture. Dosing at 2 mM or 20 mM DMG greatly shortened the 4°C TMA culture lag phase of *M. seepicolus* but did not
elevate the methane production rate (Fig. 7a), while DMG dosing did not affect the 18°C culture (Fig. S3a). Notably, DMG dosing did not increase the final methane yield of the TMA culture, indicating that DMG acts only as a cold protectant but not a methanogenic substrate and can be accumulated during GBT demethylation by Methanococcoides.

DMG not only promoted the growth rate of the O. seepicola in GBT at 4°C but also the final cell yields (Fig. 7b), likely by elevating the GBT utilization, as DMG was not used as a carbon source. Remarkably, dosing 2 mmol and 20 mmol of DMG to 4°C cocultures led to the GBT-derived methane production rate being elevated from 0.02 mmol·day⁻¹ to 0.035 mmol·day⁻¹ and 0.05 mmol·day⁻¹, respectively (Fig. 7c). In contrast, DMG dosing did not promote the GBT culture of O. seepicola (Fig. S3b) nor the coculture in methane production rates at 18°C (Fig. S3c). Consequently, these experimental results demonstrate that DMG acts as a cold protectant for both the cold seep bacteria and archaea. Therefore, the widely distributed osmoprotectant GBT could play dual roles by acting as the major precursor for methane production in addition to the cold protectant DMG in cold seep communities.

FIG 7 Dimethylglycine promotes the growth of both Methanococcoides seepicola and Oceanirhabdus seepicola in mono- and cocultures at 4°C. Mid-exponential cultures of M. seepicola and O. seepicola were inoculated at 10% into YTG basal medium containing 20 mM GBT or TMA (a) and 20 mM GBT or DMG (b), and 10% of each into GBT medium to develop a coculture (c), respectively. Dimethylglycine (DMG) at final concentrations of 2 mM and 20 mM were, respectively, dosed to the three cultures. Mono- and cocultures were grown at 4°C and 18°C (Fig. S2), respectively. Methane production (a and c) and OD₆₀₀ (b) were measured during incubation, respectively. Rates of methane production and growth were calculated from the linear portions of the curves. Triplicate cultures were assayed, and the averages and standard deviations are shown.
DISCUSSION

Methane seeps are globally widespread seafloor ecosystems wherein methane gas is abundant. Chemical studies have recently indicated that biogenic methane comprises a substantial portion of cold seep methane, like in the Sonora Margin and SCS cold seeps (11, 19). In this study, detection of methane emissions from cold seep microcosms amended with various substrates indicated that GBT was the primary methanogenic precursor. Investigation and identification of the predominant species and metabolic genes within the SCS Formosa cold seep communities and GBT-amended microcosms also indicated that GBT reduction and demethylation were predicted as the primary pathways underlying GBT-derived methane in the cold seep. The bacterium *Oceanirhabdus seepicola* and the methanogenic archaeon *Methanococcoides seepicolus* were isolated from cold seep GBT microcosms, and both represent novel species of psychrophiles that grew optimally at 18°C. The former efficiently reduced GBT to TMA via the GBT reduction pathway, while the latter converted both TMA and GBT to CH₄ via the GBT demethylation pathway. The two species were cocultured and stoichiometrically converted GBT to methane. Remarkably, the demethylated GBT product, DMG, functioned as a cryoprotectant for *O. seepicola* and *M. seepicolus* when grown at 4°C, the in situ cold seep temperature. The metagenomic and experimental data led to the proposal of a model (Fig. 8) for how the cold seep bacteria and methanogens syntrophically convert GBT to methane, which ultimately contributes to the sourcing of biogenic methane in cold seeps.

GBT is an osmoprotectant used by a wide variety of organisms, particularly those that live in oceans. Consequently, GBT is ubiquitous in coastal and marine sediments. GBT can be degraded through oxidative or reductive pathways. In oxic environments, it is demethylated by monooxygenases to yield dimethylglycine and/or sarcosine monomethylglycine and glycine (10). In anoxic environments, GBT is either reduced to TMA and acetyl phosphate or demethylated to dimethylglycine (23, 33, 34) with a methyl group being transferred to a specific methyl carrier. The selenocysteine-containing GBT reductase complex (GrdHI) is key to GBT reduction, while thioredoxin (TrxA) and its reductase (TrxB) transfer electrons from NAD(P)H to GBT reductase. In addition, *grdABCD* encodes a reductase complex that catalyzes glycine, sarcosine, or GBT reduction. A BCCT-type GBT transporter, OpuD, imports GBT to cells. The anaerobic GBT demethylation can also be accomplished by methanogenic archaea (23),...
sulfate-reducing bacteria (34), and acetogenic bacteria (35) using GBT transferase (MtgB). These anaerobes typically further metabolize the methyl group generated by GBT demethylation, and it is either transferred to coenzyme M by methanogens in methane production or through the Wood-Ljungdahl pathway (WLP) to produce acetate in bacteria (36, 37).

The genes involved in the two pathways of anaerobic GBT metabolism were all identified in metagenomic sequencing contigs derived from the Formosa cold seep sediment communities. Most of the genes involved in GBT reduction were derived from *Clostridium* and a cold seep bacterium *Psychrilyobacter* affiliated with the *Fusobacteriaceae* family. In contrast, the identified MtgB homologs were likely from additional species, mainly *Firmicutes* and *Alphaproteobacteria* (24). Similarly, genes identified that could be involved in GBT demethylation were attributed to *Chloroflexi*, *Clostridium*, and *Alphaproteobacteria*. Consequently, the two GBT metabolism pathways were simultaneously active in this cold seep.

The synergism of bacteria, particularly sulfate-reducing bacteria and methanogenic archaea in the conversion of GBT to methane, has been observed in marine ecosystems and coastal saltmarsh sediments (26, 30). Although different bacterial species implicated in GBT reduction were observed in the Formosa cold seep, the syntrophic methanogens all belonged to the *Methanococcoides* genus. Nevertheless, *Methanococcoides* strains from diverse marine habitats only demethylate some GBT (23), while *M. seepicolus* is capable of completely converting GBT to methane (Fig. 5b) and, presumably, dimethylglycine (DMG). A novel GBT-utilizing *Methanobubulus* strain was isolated from an estuary (21), which, together with *M. seepicolus*, increases the known diversity of methanogens that are capable of directly using GBT. Furthermore, the *Methanococcoides* strains that exhibit high 16S rRNA gene identities with *M. seepicolus* are widely distributed in subsea floor sediments (Table S4), indicating that GBT demethylation mediated by methanogenic archaea could be a prevalent pathway in marine ecosystems.

A 1:2.6 ratio of GBT to methane was observed in the consortium of *O. seepicola* and *M. seepicolus*, which differs from the 1:1 molecular ratio of GBT degradation to methane production previously observed in an estuarine microcosm (23). However, the ratio of 0.87 (2.6/3) methane from each methyl group is higher than the theoretical 0.75 mol methane per methyl group that is predicted from the canonical methylotrophic methanogenic pathway. Given that decreasing GBT degradation rates coincided with decreases in TMA accumulation and methane production initiation in the coculture, we assume that some of the reducing equivalent generated during GBT reduction by *O. seepicola* could be used during TMA-derived methane production. The co-occurrence of GBT degradation with TMA degradation has also been observed in estuarine microcosms (26).

Both GBT and DMG have been documented as compatible solutes in halotolerant and halophilic methanogenic archaea (38). DMG has also been observed to exhibit better heat and cold stress-relieving properties in *Bacillus subtilis*, which uses the ABC transporter OpuA to take up DMG (32). Here, evidence is provided that DMG, and presumably GBT, functions as a cryoprotectant for cold seep microorganisms, while MtgB, which is involved in DMG formation from GBT demethylation, was also encoded in the genomes of diverse cold seep bacterial species (Fig. S1b).

In conclusion, this work reports that the osmotic stress protectant, GBT, contributes to most of the biogenic methane produced in an SCS cold seep. In addition, a new bacterial species, *O. seepicola*, affiliated with the *Clostridiaceae* family, and a new methanogen species, *M. seepicolus*, which were both isolated from the cold seep, could synergistically convert GBT to methane. Moreover, *M. seepicolus* also demethylates GBT to methane and DMG, a cryoprotectant that promoted the growth of cold seep organisms at *in situ* low temperatures. GBT is an osmoprotectant ubiquitously used in marine organisms, including the cold seep benthic community, and the *grdH* genes are also found in the Gulf of Mexico cold seep. Thus,
the GBT-dependent biogenic methane production pathway identified here could be distributed among global cold seeps.

MATERIALS AND METHODS

Cold seep sediment sampling. Samples were collected from the Formosa cold seep of SCS (Fig. 1a) during the research vessel “KEKUE” scientific cruises in 2019. Reduced black sediments were collected with a gravity corer. Onboard, the top layer that is affected by seawater was removed, and the sediment column was sectioned every 20 cm, with the sediment from each section stored in anaerobic biobags at 80°C until further use.

Development of methane-producing microcosms of cold seep sediments using various methanogenic precursors. Approximately 0.5 g of cold seep sediment was inoculated into 5 ml of pre-reduced artificial seawater containing a final concentration of 20 mM TMA, GBT, acetate, choline, or 80% H2:20% CO2 gas to develop methane-producing microcosms. Each microcosm was placed in a 15-ml anaerobic tube and placed under 105 Pa of headspace gas of N2, except for the H2-CO2 treatment. A microcosm without methanogenic precursors was included as a control, and another microcosm without methanogenic precursor was amended with ampicillin and kanamycin, each at a final concentration of 100 mM to suppress bacterial growth. All the enrichments were incubated at 18°C, and methane production was monitored using gas chromatography as described below. Each experiment was conducted with four replicates.

Isolation, identification, and characterization of GBT-degrading bacteria and methanogenic archaea. Bacterial and methanogenic archaeal strains were isolated using GBT as the sole carbon source from the GBT microcosm using the Hungate rolling tube method (39). About 0.5 ml of microcosm slurry was used as the inoculant and then diluted over a 10-fold series into 5 ml prerduced YTG (40) agar medium containing 20 mM GBT. Ampicillin and kanamycin, each at a 100 mM final concentration, were added to promote isolation of methanogenic archaea. The Hungate roll tubes were incubated at 18°C, and single colonies were picked inside an anaerobic glove box (Thermo Fisher, USA).

Cell morphologies were examined with an optical microscope (BX40; Olympus). Substrate utilization by strains ZWT and LLY was assayed in YTG basal medium by omitting yeast extract, tryptone, and glucose, and with 20 mM of each test substance individually.

Growth of the bacterial strain ZWT and the methanogenic strain LLY were determined by optical density at 600 nm (OD600) measurements and methane production, respectively. Growth characteristics were assayed in YTG basal medium containing GBT. Growth temperatures were evaluated at 0, 4, 8, 18, 22, 30, and 37°C and pH-dependent growth at pH 3.0, 4.0, 5.0, 6.0, 6.5, 7.0, 7.5, 8.0, 8.5, 9.0, and 10.0. NaCl tolerance was determined at 0.0, 0.5, 1.0, 1.5, 2.0, 2.5, 3.0, 3.5, 4.0, 4.5, 5.0, 6.0, 7.0, and 7.5%. All the tests were performed in triplicate.

Measurements of methane, glycine betaine, and trimethylamine and carbon isotope analysis of CH4 and CO2. Methane was measured using a GC-14B gas chromatograph (GC; Shimadzu, Japan) equipped with a flame ionization detector and a C2 column, as previously described (41). GC temperature parameters included the column at 50°C, the injector at 80°C, and the detector at 130°C.

TMA concentration in cultures was quantified using an Agilent Technologies 6890N GC-5973N MSD equipped with a DB-Wax column (60 m × 0.25 mm [inside diameter], film thickness of 0.15 μm; J&W Scientific, Folsom, CA). Nitrogen was used as the carrier gas at a constant flow rate of 1 ml/min. The temperature of the injector was set at 280°C, the gas chromatography-mass spectrometry (GC-MS) transfer line at 280°C, the ion source at 230°C, and the quadrupole at 150°C. The reference standard solution contained 0.25 to 40 mM TMA. Spent culture (0.5 μl) was filtered through a 0.22-μm Millex GP filter (Millipore Express; polyethersulfone [PES] membrane) prior to assay measurements.

GBT was quantified using a Shim-pack G5 hydrophilic interaction liquid chromatography (HILIC) column (4.6 mm by 250 mm, 5 μm; ShimSen) that was eluted with acetonitrile/water (85:15) at a flow rate of 1.0 ml/min⁻¹ at 30°C. Spent culture liquid (200 μl) was purified using an MCI column (60 mg/3 ml; ShimSen), filtered with a 0.22-μm nylon centrifuge tube filter (Costar, Corning, NY, USA), and analyzed with a UV detector at a 195-nm wavelength. The carbon isotope signatures of methane and CO2 were determined using an isotope ratio mass spectrometer coupled with gas chromatography (GC-IRMS) instrument (Thermo Fisher Scientific, Germany) (42), with 13C/12C isotope ratios expressed as δ13C values (%) relative to the Vienna Pee Dee Belemnite (V-PDB) standard.

Illumina sequencing of archaean and bacterial 16S rRNA genes. The genomic DNAs of strains ZWT and LLY were extracted using the FastDNA Spin kit for soil (MP BioScience, Derby, UK) and quantified using a NanoDrop 2000 spectrophotometer (Thermo Fisher). The 16S rRNA genes of the cultures were amplified by PCR using the universal bacterial 16S rRNA gene amplification primers 27F (5'-AGAGTTTGATCCTGCTGCTG-3') and 1492R (5'-GGTACCTTGTGACCTT-3') and the universal methanogen 16S rRNA primers Met86F (5'-GCTACAAACAGTG-3') and Met340R (5'-CGTGTTGCTGAGG-3'), both with corresponding PCR annealing temperatures of 55°C. PCR products were ligated into T-vector plasmids, and the 16S rRNA genes were sequenced at Biosune Biotechnology Co., Ltd. (Shanghai, China). Total DNA was extracted from the in situ sediments (~0.5 g) and GBT-enriched microcosms (~0.5 ml) using a MoBio PowerSoil DNA isolation kit (MoBio, Carlsbad, CA, United States) according to the manufacturer’s protocol. The primer pairs 519F and 915R and 341F and 806R were used to amplify a 399-bp fragment of the V4-V5 hypervariable regions of archaean 16S rRNA genes (43) and a 456-bp fragment of bacterial 16S rRNA genes (44), respectively. The purified amplicons were sequenced on the
Illumina MiSeq platform at Novogene (Beijing, China) with standard protocols. Sequencing data were processed and analyzed using the QIME (version 1.7.0) pipeline (45).

Genomic and metagenomic analysis. Genomes and microcosm community DNAs were paired-end sequenced on the Illumina HiSeq 2000 platform at Novogene (Beijing, China), followed by processing of data with standard protocols. The raw paired-end metagenomic sequencing reads of two replicate samples were first quality filtered and quality controlled using the read_qc module of metaWRAP (46). Clean reads generated for each sample were individually assembled into contigs using the MEGAHIT v1.1.3 assembler (47) with default parameters, followed by removal of short contigs (\<1,000 bp). Nonredundant gene catalogs were generated as previously described (48). Briefly, open reading frames (ORFs) were predicted using MetaGeneMark (49), and those of length \( \geq 100 \) bp were retained. Redundant ORFs and contigs were removed using CD-HIT (v4.7) (50). Clean reads were then mapped to ORFs or contigs and enumerated with CoverM (https://github.com/wwood/CoverM), followed by calculation of relative abundances of genes/contigs in each sample based on the proportion of mapped reads divided by the gene/contig length. Taxonomic assignments of each ORF were generated by mapping the amino acid sequences against the NCBI nr database (https://www.ncbi.nlm.nih.gov/) using Diamond (51). Functional annotations for ORFs were predicted with eggNOG 5.0 (52) and KofamKOALA (53), and the hit score above predefined threshold of each knockout (KO) was selected.

Quantitative reverse transcription-PCR. Total RNA was extracted from mono- and coculture samples using the RNeasy Plus kit (TaKaRa, Dalian, China) according to the manufacturer’s instructions. RNA extracts were then treated with RNase-Free DNase (TaKaRa) to remove contaminating DNA, and cDNA was synthesized using the Moloney murine leukemia virus (M-MLV) reverse transcriptase cDNA synthesis kit (TaKaRa) following the manufacturer’s instructions. The transcript copies of grdH, mtgB, oupD, and mttB were quantified via qPCR using the respective primers listed in Table S5. To estimate mRNA copies of the evaluated genes, a standard curve was generated for each gene by qPCR using 10-fold serially diluted PCR products as the templates. We used 16S rRNA gene copy numbers of corresponding strains as proxies for biomass. All the measurements were performed on triplicate samples and repeated at least three times. qPCRs were performed using an SYBR Premix Ex Taq kit (TaKaRa Bio Inc., Japan) in triplicate experiments and were conducted in an ABI Prism 7000 detection system (Applied Biosystems USA). Each qPCR mixture contained 12.5 \( \mu l \) of SYBR qPCR mix (Toyobo), 5 \( \mu l \) of cDNA, 100 nM each primer, and double-distilled H\(_2\)O to a final volume of 25 \( \mu l \). PCRs were initiated at 95°C for denaturation over 30 s, followed by 35 cycles of amplification comprising 95°C at 10 s, 55°C at 30 s, and 72°C at 30 s. Fluorescence data were collected during the elongation stages.

Phylogenetic analyses. The 16S rRNA gene sequences were retrieved from the genomes of strains ZWT and LLY, and related 16S rRNA gene references were retrieved from the SILVA database for phylogenetic analysis. Phylogenetic trees were constructed using neighbor-joining methods in the MEGA 6.0 software program. Node support for both trees was evaluated by 1,000 bootstrap replicates. GrdH and MtgB phylogenies were generated from proteins encoded in metagenomic contigs for the in situ cold seep sediments and the GBT-enriched microcosms, along with references retrieved from the NCBI database. The amino acid sequences were aligned with the MUSCLE aligner, and neighbor-joining trees were constructed using the MegaX program. Node support for both trees was evaluated by 1,000 bootstrap replicates. The resulting trees were edited and visualized with iTOL (54).

Data availability. All of the data supporting the findings of this study are available in the paper and the supplemental material. The Illumina sequencing data of archaeal and bacterial 16S rRNA gene V4-V5 hypervariable regions are deposited in the NCBI Sequence Read Archive under accession numbers SRR14320062 to SRR14320068. The 16S rRNA gene sequences for amplicons from strains ZWT and LLY are deposited in NCBI GenBank under accession numbers MW674679 and MW675309, respectively. The raw metagenome sequencing data were deposited under the accession numbers SRR14270296, SRR14270297, and SRR14339844 to SRR14339846 in the NCBI Sequence Read Archive. The metagenomic contig data for the sediment and GBT enrichment are available through figshare at https://doi.org/10.6084/m9.figshare.14447793.v1.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.

FIG S1, TIF file, 1.8 MB.
FIG S2, TIF file, 0.9 MB.
FIG S3, TIF file, 0.4 MB.
TABLE S1, DOCX file, 0.01 MB.
TABLE S2, DOCX file, 0.01 MB.
TABLE S3, DOCX file, 0.01 MB.
TABLE S4, DOCX file, 0.03 MB.
TABLE S5, DOCX file, 0.02 MB.

ACKNOWLEDGMENTS
This work was funded by the National Key R&D Program of China (2018YFC0310801), the Senior User Project of RV KEXUE (KEXUE2019GZ05) and Center for Ocean Mega-Science, Chinese Academy of Sciences. We acknowledge the support of the research...
vessel KEXUE of the National Major Science and Technology Infrastructure from the Chinese Academy of Sciences in sample collection. We declare no competing financial interests.

REFERENCES

1. Levin L. 2005. Ecology of cold seep sediment, p 1–46. In Oceanography and Marine Biology, 1st ed. CRC Press, Boca Raton, FL.

2. Jorgensen BB, Boetius A. 2007. Feast and famine microbial life in the deep-sea bed. Nat Rev Microbiol 5:770–781. https://doi.org/10.1038/nrmicro1745

3. Crucq P, Vigneran A, Pignet P, Capracs JC, Lesongeur F, Toffin L, Godfrey A, Cambron-Bonavita MA. 2015. Microbial communities associated with benthic faunal assemblages at cold seep sediments of the Sonora Margin, Guayas Basin. Front Mar Sci 2:1–16. https://doi.org/10.3389/fmars.2015.00053

4. Niemann H, Løsekkann T, de Beer D, Elvert M, Nadalig T, Knittel K, Amann R, Sauer EJ, Schlüter M, Klages M, Foucher JP, Boetius A. 2006. Novel microbial communities of the Haakon Mosby mud volcano and their role as a methane sink. Nature 433:854–858. https://doi.org/10.1038/nature03227

5. Knittel K, Løsekkann T, Boetius A, Kort R, Amann R. 2005. Diversity and distribution of methanotrophic archaea at cold seeps. Appl Environ Microbiol 71:467–479. https://doi.org/10.1128/AEM.71.1.467-479.2005

6. Vigneron A, Alspø EB, Crucq P, Philibert G, King B, Baksmaty L, Lavallée D, Lomans BP, Eloe-Fadrosh E, Kyrpides NC, Head IM, Tsesmetzis N. 2019. Contrasting pathways for anaerobic methane oxidation in Gulf of Mexico cold seep sediments. mSystems 4:e00091-18. https://doi.org/10.1128/mSystems.00091-18

7. Jørgensen BB, Boetius A. 2007. Feast and famine microbial life in the deep-sea bed. Nat Rev Microbiol 5:770–781. https://doi.org/10.1038/nrmicro1745

8. Vigneron A, L’Haridon S, Godfrey A, Rousseau MG, Cragg BA, Parkes RJ, Toffin L. 2015. Evidence of active methanogen communities in shallow sediments of the Sonora Margin cold seeps. Appl Environ Microbiol 81:3451–3459. https://doi.org/10.1128/AEM.00147-15

9. Yin X, Wu W, Maek SA, Richter-Heitmann T, Kulkarni AC, Oni OE, Wendt E, Elvert M, Friedrich MW. 2019. CO2 conversion to methane and biomass in obligate methylo trophic methanogens in marine sediments. ISME J 13:2107–2119. https://doi.org/10.1038/s41396-019-0425-9

10. Wargo MJ. 2013. Homeostasis and catabolism of choline and glycine betaine: lessons from Pseudomonas aeruginosa. Appl Environ Microbiol 79:2112–2120. https://doi.org/10.1128/AEM.03565-12

11. Cao L, Lian C, Zhang X, Zhang H, Wang H, Zhou L, Wang M, Chen H, Luan Z, Li C. 2021. In situ detection of the microbial communities associated with sulfate reduction. Microbiol Mol Biol Rev 83:e00074-18. https://doi.org/10.1128/MMBR.00074-18

12. Cui H, Su X, Chen F, Holland M, Yang S, Liang J, Su P, Dong H, Hou W. 2016. Evidence of active methanogen communities in shallow sediments of the Sonora Margin cold seeps. Appl Environ Microbiol 82:289–293. https://doi.org/10.1128/AEM.03076-16

13. Feng D, Cheng M, Kiel S, Qiu J-W, Yang Q, Zhou H, Peng Y, Chen D. 2015. Dimethylglycine provides a novel carbon and energy source for methanogens (Methanococcoides spp.). Appl Environ Microbiol 81:2365–2372. https://doi.org/10.1128/AEM.03076-14

14. Zhilina TN, Zavarzin GA. 1990. Extremely halophilic, methylotrophic, anaerobic bacteria of the genus Methanococcus. FEMS Microbiol Lett 73:315–322. https://doi.org/10.1111/1574-6966.1990.tb04930.x

15. Prat L, Heinemann IU, Aerni HR, Rinehart J, O’Donoghue P, Soll D. 2012. Carbon source-dependent expansion of the genetic code in bacteria. Proc Natl Acad Sci U S A 109:21070–21075. https://doi.org/10.1073/pnas.1218613110

16. Sun J, Mauz MA, Chen Y, Giovannoni SJ. 2019. Microbial trimethylamine metabolism in marine environments. Environ Microbiol 21:513–520. https://doi.org/10.1111/1462-2920.14461

17. King GM. 1984. Metabolism of trimethylamine, choline, and glycine betaine by sulfate-reducing and methanogenic bacteria in marine sediments. Appl Environ Microbiol 48:719–725. https://doi.org/10.1128/aem.48.4.719-725.1984

18. Ye WW, Wang XL, Zhang XH, Zhang GL. 2020. Methane production inoxic seawater of the western North Pacific and its marginal seas. Limnol Oceano 65:2352–2365. https://doi.org/10.1002/lno.11457
36. Ragsdale SW. 2008. Enzymology of the wood-Ljungdahl pathway of acetogenesis. Ann N Y Acad Sci 1125:129–136. https://doi.org/10.1196/annals.1419.015.

37. Schuchmann K, Muller V. 2014. Autotrophy at the thermodynamic limit of life: a model for energy conservation in aceticogenic bacteria. Nat Rev Microbiol 12:809–821. https://doi.org/10.1038/nrmicro3365.

38. Sowers KR, Gunsalus RP. 1995. Halotolerance in Methanosarcina spp.: role of Nisup(epsilon)-acetyl-(beta)-lysine, (alpha)-glutamate, glycine betaine, and K(sup+) as compatible solutes for osmotic adaptation. Appl Environ Microbiol 61:4382–4388. https://doi.org/10.1128/aem.61.12.4382-4388.1995.

39. Hungate RE. 1969. A roll tube method for cultivation of strict anaerobes. Methods Microbiol 38:117–132. https://doi.org/10.1016/S0072-9796(08)70503-8.

40. Zhang C, Liu X, Dong X. 2004. Methanogenesis in bovine papillomatous digital dermatitis in Holstein dairy cows. J Vet Med B 51:1659–1665. https://doi.org/10.1002/jmv.2001739.

41. Zhang C, Liu X, Dong X. 2004. Syntrophomonas curvata sp. nov., an anaerobe that degrades fatty acids in co-culture with methanogens. Int J Syst Evol Microbiol 54:969–973. https://doi.org/10.1099/ijsem.0.02903-0.

42. Al G, Zhu J, Dong X, Sun T. 2013. Simultaneous characterization of methane and carbon dioxide produced by cultured methanogens using gas chromatography/isotope ratio mass spectrometry and gas chromatography/mass spectrometry. Rapid Commun Mass Spectrom 27:1935–1944. https://doi.org/10.1002/rcm.6651.

43. Santos TM, Pereira RV, Caixeta LS, Guard CL, Bicalho RC. 2012. Microbial diversity in bovine papillomatous digital dermatitis in Holstein dairy cows from upstate New York. FEMS Microbiol Ecol 79:518–529. https://doi.org/10.1111/j.1574-6941.2011.01234.x.

44. Zakrzewski M, Goernann A, Jaenicke S, Junemann S, Eikmeyer F, Szczepanowski R, Al-Soud WA, Sorensen S, Puhler A, Schluter A. 2012. Profiling of the metabolically active community from a production-scale biogas plant by means of high-throughput metatranscriptome sequencing. J Biotechnol 158:248–258. https://doi.org/10.1016/j.jbiotec.2012.01.020.

45. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, Fierer N, Peña AG, Goodrich JK, Gordon JI, Huttenhower C, Kelley ST, Knights D, Koenig JE, Ley RE, Lozupone CA, McDonald D, Muegge BD, Pirrung M, Reeder J, Sevinsky JR, Turnbaugh PJ, Walters WA, Widmann J, Yatsunenko T, Zaneveld J, Knight R. 2010. QIIME allows analysis of high-throughput community sequencing data. Nat Methods 7:335–336. https://doi.org/10.1038/nmeth.f.303.

46. Uritsky GV, DiRuggiero J, Taylor J. 2018. MetaWRAP—a flexible pipeline for genome-resolved metagenomic data analysis. Microbiome 6:158. https://doi.org/10.1186/s40168-018-0541-1.

47. Li D, Liu CM, Luo R, Sadakane K, Lam TW. 2015. MEGAHIT: an ultra-fast single-node solution for large and complex metagenomics assembly via succinct de Bruijn graph. Bioinformatics 31:1674–1676. https://doi.org/10.1093/bioinformatics/btv033.

48. Zhao D, Zhang S, Xue Q, Chen J, Zhou J, Cheng F, Li M, Zhu Y, Yu H, Hu S, Zheng Y, Liu S, Xiang H. 2020. Abundant taxa and favorable pathways in the microbiome of soda-saline lakes in inner Mongolia. Front Microbiol 11:1740. https://doi.org/10.3389/fmicb.2020.01740.

49. Zhu W, Lomsadze A, Borodovsky M. 2010. Abinitio gene identification in metagenomic sequences. Nucleic Acids Res 38:132. https://doi.org/10.1093/nar/gkq275.

50. Li W, Godzik A. 2006. Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences. Bioinformatics 22:1658–1659. https://doi.org/10.1093/bioinformatics/btl158.

51. Buchfink B, Xie C, Huson DH. 2015. Fast and sensitive protein alignment using DIAMOND. Nat Methods 12:59–66. https://doi.org/10.1038/nmeth.3176.

52. Huerta-Cepas J, Szklarczyk D, Heller D, Hernandez-Plaza A, Forslund SK, Cook H, Mende DR, Letunic I, Rattei T, Jensen LJ, von Mering C, Bork P. 2019. eggNOG 5.0: a hierarchical, functionally and phylogenetically annotated orthology resource based on 5090 organisms and 2502 viruses. Nucleic Acids Res 47:316–322. https://doi.org/10.1093/nar/gky1085.

53. Aramaki T, Blanc-Mathieu R, Endo H, Ohkubo K, Kanehisa M, Goto S, Waku K, Ogata H. 2020. KofamKOALA: KEGG ortholog assignment based on profile HMM and adaptive score threshold. Bioinformatics 36:2251–2252. https://doi.org/10.1093/bioinformatics/btz859.

54. Letunic I, Bork P. 2019. Interactive Tree Of Life (ITOL) v4: recent updates and new developments. Nucleic Acids Res 47:W256–W259. https://doi.org/10.1093/nar/gkz239.