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Genomic and Transcriptomic Evidence Supports Methane Metabolism in *Archaeoglobi*

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ABSTRACT Euryarchaeal lineages have been believed to have a methanogenic last common ancestor. However, members of euryarchaeal *Archaeoglobi* have long been considered nonmethanogenic and their evolutionary history remains elusive. Here, three high-quality metagenomic-assembled genomes (MAGs) retrieved from high-temperature oil reservoir and hot springs, together with three newly assembled *Archaeoglobi* MAGs from previously reported hot spring metagenomes, are demonstrated to represent a novel genus of *Archaeoglobaceae*, *Candidatus Methanomixophus.* All *Ca. Methanomixophus* MAGs encode an M methyltransferase (MTR) complex and a traditional type of methyl-coenzyme M reductase (MCR) complex, which is different from the divergent MCR complexes found in *Ca. Polytropus marinifundus.* In addition, *Ca. Methanomixophus dualitatem* MAGs preserve the genomic capacity for dissimilatory sulfate reduction. Comparative phylogenetic analysis supports a laterally transferred origin for an MCR complex and vertical heritage of the MTR complex in this lineage. Metatranscriptomic analysis revealed concomitant in situ activity of hydrogen-dependent methylotrophic methanogenesis and heterotrophic fermentation within populations of *Ca. Methanomixophus hydrogenotrophicum* in a high-temperature oil reservoir.

IMPORTANCE Current understanding of the diversity, biology, and ecology of *Archaeae* is very limited, especially considering how few of the known phyla have been cultured or genomically explored. The reconstruction of *Ca. Methanomixophus* MAGs not only expands the known range of metabolic versatility of the members of *Archaeoglobi* but also suggests that the phylogenetic distribution of MCR and MTR complexes is even wider than previously anticipated.

KEYWORDS horizontal gene transfer, HGT, metatranscriptomics, methyl-coenzyme M reductase complex, MCR complex, oil reservoir, methanogens

Methanogenic lifestyles were originally inferred to be restricted to a few “traditional” *Euryarchaeota* that were distributed within seven orders, including *Methanopyrales, Methanococcales, Methanobacterales, Methanosarcinales, Methanocellales, Methanomicrobiales,* and *Methanomassiliicoccales* (1, 2). The recent discovery of Methanotronarchaeia, a novel extreme halophilic methanogen which branches basally to *Halobacteria,* provides new insights into the evolutionary transition from an anaerobic and methanogenic ancestor to haloarchaeal lineages (3, 4). In addition, culture-independent approaches revealed a growing number of archaeal groups outside the
Euryarchaeota (“Ca. Bathyarchaeota,” “Ca. Verstraetearchaeota,” and “Ca. Helarchaeota”) whose genomes contain the key genes for methanogenesis (5–9), suggesting the evolutionary origin of methanogenesis predates the Euryarchaeota.

As key enzymes in archaeal methane cycling, methyl-coenzyme M reductase (MCR) complexes can be divided into two main clusters, with one cluster being composed of MCR complexes from traditional euryarchaeal lineages and “Ca. Verstraetearchaeota.” MCR complexes in this cluster catalyze the final step of methanogenesis or the initial step in anaerobic methane oxidation (1). On the other hand, the MCR homologs found in “Ca. Bathyarchaeota” formed another deep divergent branch, together with a newly found MCR complex which catalyzed short-chain alkane (butane and, probably, propane) in a euryarchaeal lineage, “Ca. Syntrophoarchaeum” (10). The close association of MCR complexes and the shared metabolic features of the “Ca. Bathyarchaeota” and “Ca. Syntrophoarchaeum” suggest that the MCR complex in “Ca. Bathyarchaeota” may catalyze short-chain hydrocarbon oxidation rather than methane production (10).

Despite being phylogenetically close to methanogenic Euryarchaeota, members of Archaeoglobi have long been considered nonmethanogenic archaea (11, 12). Genes conserved in both hydrogenotrophic methanogenesis and archaeal type Wood-Ljungdahl (WL) pathway are present in Archaeoglobi genomes, suggesting the remnants of its ancestral methane-cycling lifestyle (13). For a long time, however, genes encoding methyl coenzyme M reductase complex (MCR) and a complete N 5-methyl-H₄M(S)PT:coenzyme M methyltransferase (MTR) complex had not been found in representatives of Archaeoglobi, which are hypothesized to have been lost during evolution after receiving dsrAB genes from the bacterial members via horizontal gene transfer (HGT) (13, 14). The MTR complex, encoded by the mtr operon, catalyzes the energy-conserving (sodium-pumping) methyl transfer from H₄M(S)PT to CoM (15), which is one of the key enzymes in hydrogenotrophic methanogenesis (16). Recently, Boyd et al. found two divergent MCR complexes in a metagenome-assembled genome (MAG) representing a basal member of the class Archaeoglobi, “Ca. Polytropus marinifundus,” which could utilize nitrate, iron, and sulfur compounds as electron acceptors (17). However, genes coding for MTR complex were largely missing in “Ca. Polytropus marinifundus” except for mtrH, ruling out the possibility of conserving energy from hydrogenotrophic methanogenesis (17). Further analysis revealed that the two divergent MCRs were most likely received from “Ca. Syntrophoarchaeum” and “Ca. Bathyarchaeota” via HGT, which suggests their potential role in hydrocarbon activation (17). Hence, data explaining what the lowest common ancestor (LCA) of Archaeoglobi looks like and how the evolution transition occurs is still elusive.

Here, we assembled three Archaeoglobi MAGs and collected another three newly assembled Archaeoglobi MAGs from a previous study (7). All these MAGs were retrieved from samples from subsurface hydrothermal environments, such as hot springs and oil reservoir, indicating an anoxic and thermophilic life style of this new lineage. The comparative genomic analysis of these six MAGs expanded the current knowledge about the evolution trend of Archaeoglobi members (17).

RESULTS AND DISCUSSION

Discovery of a novel Archaeoglobaceae genus, “Ca. Methanomixophus.” In a previous study, microbial biomass from formation waters was collected from the Jiangsu oil reservoir (18). The combined metagenomes were coassembled, and the resulting contigs were binned into 44 unique genomes (18). As revealed by the genome tree, one of the high-quality MAGs (Bin16) was phylogenetically placed into the Archaeoglobi clade (Fig. 1). However, the annotation of Bin16 demonstrated a distinct genotype of Archaeoglobus species. Surprisingly, nearly the whole set of genes associated with reversible hydrogenotrophic methanogenesis was found in Bin16, including a methyl–coenzyme M reductase (McrABG) complex (Fig. 1) and a methyl-H₄M(S)PT: coenzyme M methyltransferase complex (MtrABCDEFGH) (Fig. 1; see also Table S4 in the supplemental material). Further phylogenetic analysis based on the concatenated amino acid alignment of McrABG placed Bin16 close to the traditional McrABGs rather
FIG 1 Phylogenetic tree showing the placement of "Ca. Methanomixophus" and the distribution of studied genes. The tree was inferred from the concatenation of 400 conserved marker genes using PhyloPhlAn (65), and only lineages containing MCR complex-bearing genomes for gene distribution are exhibited, whereas the lineages without MCR complex detected so far are outlined with dashed lines. Colored and blank squares represent the presence and absence of annotated genes, respectively. Genes associated with nitrate reduction and sulfate reduction are marked in brown and red, respectively; genes for iron reduction are marked in dark blue; genes involved in methanogenesis are marked in light blue; genes for beta-oxidation are marked in pink; genes associated with biosynthesis are marked in green. A functional complex consisting of multiple subunits was considered present if ≥75% of the genes comprising the complex were detected. Branches that represent "Ca. Methanomixophus" are marked in red, and the clade that represents "Ca. Polytropus marinifundus" is marked in blue. Abbreviations are as follows: nar, nitrate reductase/nitrite oxidoreductase; nap, periplasmic nitrate reductase; nir, nitrite reductase (NO-forming); nor, nitric oxide reductase; sat, sulfate adenylyltransferase; apr, adenylylsulfate reductase; dsr, dissimilatory sulfite reductase; qmo, quinone-modifying oxidoreductase; dsrMKJOP, membrane-bound heterodisulfide reductase; fwd, formylmethanofuran dehydrogenase; frh, coenzyme F420 hydrogenase; ehb, energy-converting hydrogenase A/B/C; mch, multiheme c-type cytochromes; fadA, acyl-CoAacyltransferase; aco, acyl-CoA acyltransferase; FadD, long-chain acyl-CoA synthetase; acoa/adcd/bcd, acyl-ACP dehydrogenase; palA/crt/ish/fadB/ech, enoyl-CoA hydratase; fadA/fad/ish/ech/paol, 3-hydroxyacyl-CoA dehydrogenase; MCR, multiheme c-type cytochromes; fadA, acyl-CoA acyltransferase; atoB, acyl-CoA C-acyltransferase; Mnh, genes for biosynthesis of the menaquinone; Mpt, genes for biosynthesis of the methanophenazine.
than to the divergent cluster (Fig. 2C). In order to study the distribution of the new *Archaeglobi* members in nature, the *mcrA* gene in Bin16 was used to screen metagenomes in IMG publicly available (for details, see Text S1 in the supplemental material), and the closely related *mcrA* genes were detected in two thermal aquatic metagenomes: an *in situ* cellulolytic enrichment in Great Boiling Spring (Integrated Microbial Genomes identifier [IMG-ID]: 3300000106, NV, USA) (19) and a water sample from Washburn Spring (IMG-ID: 3300005860, Yellowstone National Park, USA). These metagenomes were individually assembled and differentially binned, and then two additional *Archaeglobi* MAGs, Bin11 and Bin74, were retrieved from metagenomes of Great Boiling Spring (IMG-ID: 3300000106, NV, USA) and Washburn Spring (IMG-ID: 3300005860, Yellowstone National Park, USA)
3300005860, Yellowstone National Park, USA), respectively. Similarly to Bin16, Bin11 and Bin74 also contained MCR and MTR complexes. The completeness, contamination (redundancy), and number of total contigs meet the requirements for being ranked as nearly complete genomes as proposed previously by Parks et al. (20) and as high-quality draft genomes as proposed by Bowers et al. (21) (Table 1). Subsequently, examination of the contigs containing methanogenesis-related genes in these MAGs revealed that they have sequence composition characteristics (average GC content, sequencing coverage, and tetranucleotide frequencies) typical of their respective genomes (see Fig. S1 at https://figshare.com/articles/Fig_S1_Evaluation_of_the_statistical_properties_of_scaffolds_of_Ca_M_hydrogenus_Bin16_a_Bin11_b_and_Bin74_c/9918200). To make a comprehensive study of Archaeoglobi MAGs, three newly assembled Archaeoglobi MAGs (LMO1, LMO2, and LMO3), which also contained MCR and MTR complexes homologous to those of Bin16 (Fig. 1; see also Fig. 2A and C), were downloaded from the NCBI database and included into this study (7).

Phylogenetic analysis of taxonomic markers from Bin16, Bin11, Bin74, and LMO1 to LMO3 placed their position close to the Archaeoglobus fulgidus lineage, forming two separate clusters (Fig. 1; see also Table S8). These MAGs shared 71% to 74% of orthologous genes with the A. fulgidus genome (see Fig. S2 at https://figshare.com/articles/Fig_S2_Pairwise_comparison_of_shared_orthologous_gene_sequences_across_genomes_of_Archaeoglobi_isolates_and_Ca_Methanomixophus_MAGs_/9918206), and the average amino acid identities (AAI) between these MAGs and other Archaeoglobus proteomes were found to be below 65% (see Fig. S2 at https://figshare.com/articles/Fig_S2_Pairwise_comparison_of_shared_orthologous_gene_sequences_across_genomes_of_Archaeoglobi_isolates_and_Ca_Methanomixophus_MAGs_/9918206), hinting at a novel Archaeoglobaceae genus according to the category thresholds proposed by Konstantinidis et al. (22). The proposal for a novel genus was supported by analyses performed with the GTDBtk tool (https://github.com/Ecogenomics/GTDBTk), which uses a recently described relative evolutionary distance metric to predict the divergence of newly binned clades (23) (Table S5). The 16S rRNA gene fragments found in Bin11 (370 bp), Bin74 (918 bp), LMO1 (1,213 bp), and LMO3 (356 bp) showed 92% to 93% similarity to A. fulgidus DSM 4304 (GenBank accession number AE000782.1), which also suggested a novel genus-level lineage of Archaeoglobaceae for this clade (24). Phylogenetic analysis of these 16S rRNA gene sequences placed them into a monophyletic cluster with other uncultured Archaeoglobus clone sequences from similar environments from which these MAGs have been retrieved, such as oil reservoirs (GenBank accession numbers GU179414, KY707708, and JN794070) and hot springs (GenBank accession numbers EU924223 and KP784730); however, the results also showed that they were distantly related to A. fulgidus sequence clusters (Fig. 3). Consistent with the genome tree, 16S rRNA gene sequences from Bin11 and Bin74 were placed into a cluster separate from that containing LMO1 and LMO3 (Fig. 3). On the basis of these results, we proposed the name “Ca. Methanomixophus hydrogenotrophicum” gen. nov., sp. nov., for Bin11, Bin16, Bin74, and LMO2, in which “Methano” stands for methane, “mixo” means mixotrophy, and “hy-

### Table 1: Summary statistics of new “Ca. Methanomixophus” MAGs

| MAG   | Completeness (%) | Redundancy (%) | No. of contigs | Total length (bp) | GC content (%) | No. of CDS | No. of RNAs | Relative abundance (%) | Relative activity (%) | RAST-ID or reference |
|-------|------------------|----------------|----------------|-------------------|----------------|------------|-------------|------------------------|----------------------|---------------------|
| Bin16 | 96.5             | 1.3            | 45             | 1,661,519         | 45.5           | 1,834      | 46          | 1.51                   | 0.06                 | 6666666.39          |
| Bin11 | 94.6             | 1.6            | 133            | 1,403,156         | 47.6           | 1,389      | 31          | 3.05                   | 0.27                 | 6666666.23          |
| Bin74 | 96.1             | 1.3            | 126            | 1,532,438         | 45.6           | 1,895      | 39          | 3.05                   | 0.27                 | 6666666.39          |
| LMO1  | 88.89            | 1.31           | 140            | 1,557,307         | 43.8           | 1,851      | 46          | 7                      | 7                    | 6                 |
| LMO2  | 88.03            | 1.96           | 220            | 1,514,415         | 45.9           | 1,900      | 45          | 7                      | 7                    | 6                 |
| LMO3  | 97.6             | 0              | 135            | 1,567,523         | 43.9           | 1,872      | 37          | 7                      | 7                    | 6                 |

*Data were calculated by mapping quality-controlled metagenome/metatranscriptome short reads to the MAG nucleotide files using Bowtie2 (69) and were filtered for MapQ values of >2.*
drogenotrophicum” indicates the potential capability of utilizing hydrogen molecules in this lineage, and the “Ca. Methanomixophus dualitatem” sp. nov. for LMO1 and LMO3, in which “dualitatem” indicates the uncertain energy conservation strategy utilized in this lineage, considering that the genomic capacities of both anaerobic methane metabolism and dissimilatory sulfur metabolism were preserved (see below).

Evolutionary history of “Ca. Methanomixophus” MCR complex and MTR complex. An operon (mtrABCDEFGH) consisting of genes that encode the complete methyl-H₄M(S)PT:coenzyme M methyltransferase complex was found in all “Ca. Methanomixophus” MAGs (Fig. 1; see also Table S4). The blastp search revealed that these mtrABCDEFGH genes were predominantly associated with methanogens at 38% to 71% amino acid identity (Fig. 4A), while genes located upstream and downstream in the same contigs were found to be mostly related to *Archaeoglobi* species with average amino acid identity of 63% (Fig. 4A). Metagenomic short reads were mapped to these contigs, and the well-overlapped alignments confirmed the assembly (Fig. 4B). Furthermore, the assembly of the contig in “Ca. Methanomixophus hydrogenotrophicum” Bin16 was checked by PCR amplifications using DNA of W2 and W9 production water samples from the Jiangsu oil reservoir (18), and the amplicon sequences matched 99% to 100% to the partial *mtrH* gene and multiple genes located downstream in the contig.

**FIG 3** Consensus tree showing the placement of the 16S rRNA genes. “Ca. Methanomixophus” 16S rRNA gene sequences are marked in red, and the “Ca. Polytopus marinifundus” 16S rRNA gene sequence is marked in blue. The cluster contains “Ca. Methanomixophus hydrogenotrophicum” and “Ca. Methanomixophus dualitatem” sequences and is shaded in cyan and pink, respectively.
The emergence of colocated genes homologous to distinct microorganisms could be a result of HGT or, as suggested, a remnant character of the LCA before gene losses. To explore the evolutionary history of the MTR complex, the gene phylogeny of the MTR complex was compared with the genome phylogeny. To get a finer phylogenetic resolution of MCR and MTR complex, concatenated amino acid alignments of subunits for the same enzyme were chosen instead of individual protein sequence since these subunits were constitutive conserved in all MCR/MTR-bearing genomes and located in the same operon (25). Consistently, the genome tree and the mtrABCDEH gene tree both resulted in a monophyletic clade of "Ca. Methanomixophus," and the branching order of "Ca. Methanomixophus" clade in mtrABCDEH gene tree is congruent with the genome tree, which branches off between Class I (Methanopyrales, Methanococcales and Methanobacteriales) and Class II (Methanomicrobia, comprised of Methanosarcinales, Methanocellales and Methanomicrobiales) methanogens (13) with strong bootstrap supports (Fig. 2A and B). No potential mobile genetic elements (such as integrons, transposons, repeat sequences and tRNAs), which are hallmarks for HGT, could be identified in the flanking regions of mtr genes (Fig. 4A; see also Table S6). Further analysis of GC content and 4mer frequencies of these mtr operons also showed consistent profiles with the surrounding gene context in the contigs (see Fig. S3 at https://figshare.com/articles/FIG_S3_Comparison_of_tetranucleotide_frequencies_and_GC_content_between_MCR_MTR_operons_and_surrounding_gene_context_in_the_scaffolds_/9918209), suggesting that either these mtr operons are inherited vertically or the acquirement of this operon through HGT did not occur in recent evolutionary history (17). These evidence collectively suggests a vertical inheritance of evolutionary history for “Ca. Methanomixophus” MTR.

(Fig. 4C; see Text S2).
Genes (*mcrABG*) encoding all subunits of methyl-coenzyme M reductase complex, where ligand-binding sites for CoB, CoM, and cofactor F₄₃₀ are well conserved (see Fig. S4 at https://figshare.com/articles/Fig_S4_Conservation_of_functionally_important_McrA_residues_including_ligand_cavity_sites_and_F420_CoB_and_CoM_binding_sites_as_revealed_by_Ermler_et_al_21_/9918236), was found in these MAGs. As revealed by structure prediction, the McrA, McrB, and McrG within these genomes showed high similarity to the crystal structures from *Methanopyrus kandleri*. (see Fig. S5 at https://figshare.com/articles/Fig_S5_Structural_modelling_and_active_sites_of_the_MCR_complex_in_Ca_M_Bin16_/9918233). BLASTP search results demonstrated that the *mcrABG* genes in these MAGs are most similar to genes in “Ca. Methanodesulfokores washburnensis” (77% to 83% amino acid identity), which is a newly assembled MAG affiliated with the Korarchaeota phylum (26). Phylogenetic analysis of the concatenated McrABG sequences also placed them with “Ca. Methanodesulfokores washburnensis” (denoted “Korarchaeota” in Fig. 2C), forming a basal branch to euryarchaeal lineages (Fig. 2C), which is incongruent with the genome tree (Fig. 2B). However, it should be noted that despite the largely congruent branching order in the MCR tree and genome tree for most members, incongruent positions were also observed for some traditional methanogens in this study (Fig. 2B and C) and in several other studies (16, 17). Further analysis revealed that the *mcr* operons in this lineage contain two extra genes, *mcrC* and *mcrD*, with unknown function (27). This arrangement of *mcrBDCGA* in an operon resembles the *mcr* operons in most conventional euryarchaeal methanogens and “Ca. Methanodesulfokores washburnensis,” rather than the arrangement of *mcrBGA* in “Ca. Verstraetearchaeota” (6) or the divergent type *mcrBAG* in some “Ca. Syntrophoarchaeum” and “Ca. Polytropus marinifundus” isolates (Fig. 2C) (17). As a result, the *mcr* operon in “Ca. Methanomixophus” genomes is likely to be accepted through HGT but would likely be from a donor different from “Ca. Polytropus marinifundus” (17) (Fig. 2C), despite the fact that we did not find any mobile elements or divergent GC or 4mer profiles surrounding the gene context of the MCR operons (Table S6; see also Fig. S3 at https://figshare.com/articles/Fig_S3_Comparison_of_tetranucleotide_frequencies_and_GC_content_between_MCR_MTR_operons_and_surrounding_gene_context_in_the_scaffolds_/9918209).

Metabolic potential and *in situ* activity of “Ca. Methanomixophus.” In contrast to all *Archaeoglobi* known so far (11), including “Ca. Polytropus marinifundus,” genes encoding key enzymes associated with dissimilatory sulfate reduction, namely, the genes encoding sulfate adenylyltransferase (*sat*), adenylylsulfate reductase (*aprAB*), and dissimilatory sulfide reductase (*dsrAB*), are absent in “Ca. Methanomixophus hydronenotrophicum” genomes (Fig. 1; see also Table S4). Further examination of metagenome contigs and short reads did not reveal any related sequences for these missing genes (see Text S1). To confirm that the loss of sulfate-reducing genes in these MAGs was not an artifact caused by an assembly mistake, we took the genes that have been found to locate upstream or downstream of sulfate-reducing genes in public *Archaeoglobus fulgidus* reference genomes (Table S2) as anchors, and the absence of genes encoding sulfate-reducing products in “Ca. Methanomixophus hydronenotrophicum” Bin16 was validated by the presence of amplicons spanning those anchor genes and their adjacent genes (see Fig. S6 at https://figshare.com/articles/Fig_S6_Contigs_containing_sulfate-reducing_genes_in_Ca_Methanomixophus_MAGs_/9918203; see also Table S3). The only exception was the presence of DsrC, encoded by the *dsrC* gene, with two conserved cysteines in the C-terminal region (see Fig. S7 at https://figshare.com/articles/Fig_S7_Trimmed_alignment_of_DsrC_TsuE_AsPA_amino_acid_sequences_/9918230). DsrC is a small protein functioning in the terminal step of sulfate reduction (28). However, without the dissimilatory sulfate reductase (*dsrAB*) gene, its role in this lineage remains unclear. In a previous study, *dsrC* genes were found in organisms that do not have the *dsrAB* genes (29) and their products were predicted to function like TusE proteins, which were shown to participate in a sulfur-relay system (30).
“Ca. Methanomixophus hydrogenotrophicum” genomes also lack genes encoding the quinone-interacting membrane-bound oxidoreductase (QmoABC) complex. The QmoABC complex was previously proposed to link the electron transfer chain to the first reductive step of sulfate reduction in *Archaeoglobus* (31). The absence of *qmoABC* genes is consistent with the absence of the aforementioned dissimilatory sulfate reduction-related genes, indicating the inability of this clade to generate ATP through sulfate reduction. Furthermore, genes for reducing other electron acceptors, such as nitrate, thiosulfate, and iron, all of which have been shown to be widely utilized by other *Archaeoglobi* species, were all missing in this clade (Fig. 1). Therefore, this new lineage might not gain energy through respiration (Fig. 5).

“Ca. Methanomixophus hydrogenotrophicum” genomes encodes a nearly complete archaeal type Wood-Ljungdahl pathway, except for the gene encoding N$_5$N$_{10}$-methylene-H$_4$MPT reductase (Mer) (Table S4). The methylene-tetrahydrofolate reductase (MetF), which was previously suggested to act as a substitute for Mer in the reverse process of methanogenesis (32), was also missing in all “Ca. Methanomixophus hy-
drogenotrophicum" genomes (Fig. 1). Given the absence of *mer* and *metF* in both genomes and unassembled metagenomic contigs, a bypass pathway for methanol-utilizing methanogenesis in which methanol is oxidized to formaldehyde and subsequently to N₅,N₁₀-methylene-H₄MPT has been proposed (32, 33) (Fig. 5). The presence of genes encoding a fusion protein of formaldehyde-activating enzyme and hexulose-6-phosphate synthase (FaeB-Hps) and encoding short-chain alcohol dehydrogenases (Aldh) and CoM methyltransferase (Mta) suggested the presence of this bypass pathway in this new clade (33, 34).

"Ca. Methanomixophus hydrogenotrophicum" genomes encode an F₄₂₀-nonreducing hydrogenase (MvhADG) (Fig. 1). In hydrogenotrophic methanogens, this hydrogenase forms a complex with heterodisulfide reductase (HdrABC) which bifurcates electrons from H₂ to reduce ferredoxin and heterodisulfide, likely coenzyme M (CoM) and coenzyme B (CoB), in methanogens and a gamma subunit of dissimilatory sulfate reductase (DsrC) in *Archaeoglobus* (35, 36). Only an HdrA subunit which contains flavin adenine dinucleotide (FAD) is present in the genomes, indicating the ability to reduce ferredoxin. However, no subunit of HdrB, which catalyzes the direct reduction of disulfide, was found. Instead, the genomes contain another gene encoding HdrD, which is a fusion of HdrB and HdrC (37), suggesting that HdrD may replace HdrB in the Mvh/Hdr complex (26) (Fig. 5). Noteworthy is the fact that the lack of an HdrB subunit has also been observed in an uncultured *Archaea* genome (26) and a sulfate-reducing bacteria (38), hinting at an underrepresented energy-conserving mechanism within anaerobic microorganisms. However, future studies on the proteome of "Ca. Methanomixophus hydrogenus" and *in vitro* investigations will be required to elucidate the role of the HdrD detected here. A gene cluster encoding an F₄₂₀H₂:quinone oxidoreductase (Fqo) was found in this clade (Fig. 1; see also Table S4). The FqoF subunit is usually considered bound to the membrane integral module of FqoBCDI, forming the Fqo complex which oxidizes F₄₂₀H₂ coupled with menaquinone (MQ) reduction in *Archaeoglobus* (39). Nevertheless, it has been proposed that the FqoF subunit might be soluble in cytoplasm and might independently catalyze the reduction of F₄₂₀ with Fd_red oxidation in *Archaeoglobus* species (40). A similar mechanism has also been proposed for the homologous Fqo in the *Methanomixophus* genomes analyzed in this study, which also contain FAD and iron-sulfur clusters, that is, mediation of the electron transfer from Fd_red onto F₄₂₀ (Fig. 5). Furthermore, as shown in another study, the Fpo/Fqo complex is capable of catalyzing a Fd:MP/Fd:MQ oxidoreductase reaction independently of the Fpo/FqoF subunit (41) (Fig. 5). Additionally, there is also a potential of the FqoF-associated Fqo complex to link Fd_red oxidation to both F₄₂₀ reduction and menaquinone reduction by flavin-dependent electron bifurcation (42) (Fig. 5). In "Ca. Methanomixophus hydrogenotrophicum," the electrons carried by menaquinone (MQ) might be then transferred through the membrane-bound heterodisulfide reductase (HdrDE) to reduce CoM-S-S-CoB (43, 44) (Fig. 5).

However, the key enzyme of hydrogenotrophic methanogenesis, F₄₂₀-reducing hydrogenase, is missing in all MAGs, which rules out the possibility of hydrogenotrophic CO₂-reducing methanogenesis in these organisms (1). On the other hand, the detection of methyltransferase (MtaABC) and Mvh:Hdr complex indicates a potential hydrogen-dependent methylotrophic (methanol) methanogenesis lifestyle in this clade, which is similar in mechanism to those seen with the members of *Methanomassilicoccales* (43) (Fig. 5). In this case, electrons from H₂ could be transferred to reduction of CoM-S-S-CoB facilitated by Fqo and Hdr complexes as mentioned above. The methyl branch of the Wood-Ljungdahl pathway and the MTR complex would function in anabolic production of acetyl coenzyme A (acetyl-CoA) from CO₂ and CH₂-CoM, respectively (Fig. 5).

The genomic features collectively suggest a methanogenic lifestyle for "Ca. Methanomixophus hydrogenotrophicum"; however, its potential role as an anaerobic methane oxidizer could not be ruled out. In such a case, "Ca. Methanomixophus hydrogenotrophicum" may oxidize methane anaerobically through a reversible methanogenesis pathway and would require syntrophic partners as electron sinks.
likely the sulfate-reducing microorganisms detected in the same environments (18, 41) (Fig. 5). And the membrane-associated electron-transfer HdrDE complexes would mediate the electron flow in a reverse direction of methanogenesis (Fig. 5).

“Ca. Methanomixophus dualitatem” genomes displayed high similarity in genomic features with “Ca. Methanomixophus hydrogenotrophicum,” containing mtr, mcr, and Mvh/Hdr complexes (Fig. 1). Interestingly, the gene coding for N5,N10-methylene-H4MPT reductase (mer) was found in “Ca. Methanomixophus dualitatem” genomes, completing the WL pathway in this lineage (Fig. 5). Hence, “Ca. Methanomixophus dualitatem” may also conserve energy through hydrogenotrophic methanogenesis (Fig. 5). “Ca. Methanomixophus dualitatem” genomes possess extra genes (sat, AprAB, dsrABC, and qmoABC) for sulfate reduction, suggesting that “Ca. Methanomixophus dualitatem” members might alternatively perform sulfate-dependent anaerobic methane oxidation in single cells (45) (Fig. 5). Interestingly, a newly assembled Korarchaeota genome in “Ca. Methanodesulfokores washburnensis” carries genes that code for a partial dissimilatory sulfate reduction pathway, as well as the co-occurring genes coding for anaerobic methane oxidation, hinting at a close link between anaerobic methane metabolism and dissimilatory sulfur reduction in Archaea members (26).

In addition, genes involved in fatty acid degradation (β-oxidation) and in degradation of peptides/amino acids, d-lactate metabolism, and acetogenesis were found in all “Ca. Methanomixophus” MAGs (Fig. 5; see also Table S4) (see Text S1 for more details), which is consistent with results from the closely related Archaeoglobus species (11).

Metatranscriptomic analysis of “Ca. Methanomixophus hydrogenotrophicum” Bin16 in sample W15 obtained from the Jiangsu oil reservoir demonstrated that most genes associated with the common pathway of methanogenesis, namely, mcrABCDEFG, mtrA-H, fwdABDEFG, and mtrD, were transcribed to different degrees (Fig. 5; see also Table S7). Unexpectedly, the transcriptional levels of individual genes coding for the subunits of Mr were found to be greatly uneven (Table S7). It is also noteworthy that the fragments per kilobase per million (FPKM) values of mcrABG genes were relatively low compared with those seen with other genes described here. Since these FPKM values were calculated based on the metatranscriptome data set representing a very complex microbial community, future research, likely performed on the basis of pure culture isolation, will be required to resolve these issues. Meanwhile, genes specific to methanogenesis from both methanol (mtaABC) and hydrogen (mvhADG) were transcribed (Fig. 5; see also Table S7), indicating an active hydrogen-dependent methylo trophic methanogenesis process of “Ca. Methanomixophus hydrogenotrophicum” in the oil reservoir. Transcripts of multiple genes involved in d-lactate degradation (dld) and in degradation of fatty acids and peptides were detected, as well as transcripts of those involved in degradation of acetyl-CoA synthetase (ADP-forming) (acd), aldehyde dehydrogenases (aldh), and alcohol dehydrogenases (adhP), suggesting that heterotrophic fermentation was also active in “Ca. Methanomixophus hydrogenotrophicum” Bin16 (Fig. 5).

Evolutionary origin of Archaeoglobus. The discovery of hyperthermophilic “Ca. Methanomixophus” sheds light on how the transition from a methanogenic LCA to nonmethanogenic Archaeoglobus, which was likely facilitated by multiple individual gene gains (through HGT) and losses, might have occurred (Fig. 6). Previous studies have shown that the dsrAB genes in Archaeoglobus species were accepted from a bacterial donor through HGT (see Fig. S8 at https://figshare.com/articles/Fig_58_Phylogenetic_trees_showing_the_placement_of_concatenated_DsrAB_sequences_/9918215) (14), and phylogenetic analysis of sat and aprAB in Archaeoglobus species and of narGHI in Ferroglobus placidus and “Ca. Polytropus marinifundus” also suggested their bacterial origin (see Fig. S9, S10, and S11 at https://figshare.com/articles/Fig_59_Phylogenetic_trees_showing_the_placement_of_concatenated_AprAB_sequences_/9918212, https://figshare.com/articles/Fig_510_Phylogenetic_trees_showing_the_placement_of_concatenated_NarGHI_sequences_/9918221, and https://figshare.com/articles/Fig_511_Phylogenetic_trees_showing_the_placement_of_the_Sat_sequences_/9918227, respectively) (14, 46). Furthermore, phylogenetic analysis
demonstrated that the sat and aprAB genes in Geoglobus species, Ferroglobus placidus, and “Ca. Polytropus marinifundus” were located in separate clusters, distantly related to Archaeoglobus species (see Fig. S9 and S11 at https://figshare.com/articles/Fig_S9_Phylogenetic_trees_showing_the_placement_of_concatenated_AprAB_sequences_/9918212 and https://figshare.com/articles/Fig_S11_Phylogenetic_trees_showing_the_placement_of_the_Sat_sequences_/9918227, respectively), which indicates multiple HGT events and different donors of these genes during evolution (Fig. 6). Consistently, the contigs in “Ca. Methanomixophus dualitatem” genomes that flank sat and apr genes showed many hallmarks of HGT (see Fig. S6 at https://figshare.com/articles/Fig_S6_Contigs_containing_sulfate-reducing_genes_in_Ca_Methanomixophus_MAGs_/9918203), including repeat regions, transposons, and tRNA genes often found in association with genomic islands (47), and are common target sites for phage attach-ment and integration (48). Overall, sulfate-reducing ability was largely retained in Archaeoglobus members as well as in “Ca. Methanomixophus dualitatem,” as found in this study, which usually inhabit submarine hydrothermal environments (49, 50), hot springs (51), and deep oil reservoirs (52), with elevated concentrations of carbon dioxide, methane, hydrogen, hydrogen sulfide, and sulfate.

The recent discovered hydrogenotrophic methanogenesis pathway in “Ca. Verstratenarchaeota,” which had previously been considered a genus of strict methylotrophic methanogens, implies an ancient origin of hydrogenotrophic methanogenesis and a later adaptation of methylotrophic methanogenesis for all methanogens (53). On the basis of the detection of a MTR complex and a traditional MCR complex in this novel Archaeoglobi genus, as well as the widespread WL pathway within Archaeoglobi lineages, we propose a similar evolutionary trend for Archaeoglobi: hydrogenotrophic methanogenesis first evolved to support life in a nutrient-poor environment that required using only CO2 as a source of carbon, and then complex carbon source...
degradation ability evolved as more-complex nutrient environments became available, followed by accepting respiring ability using different electron sinks from bacterial members, which made them more metabolically flexible under different environmental conditions (Fig. 6). The previous discovery of the basal member of Archaeoglobi, “Ca. Polytropus marinifundus,” suggests the acquisition of the divergent MCR complex through a HGT event (17). In this study, however, we expanded the current knowledge by showing that (i) two separate HGT events of different types of MCR complexes, from different donors, may have occurred (Fig. 6) and that (ii) despite the fact that the original MCR and MTR complexes encoded in the Archaeoglobi LCA have been substantially lost in most Archaeoglobi lineages, the clade of “Ca. Methanomixophus” retains the original MTR complex (Fig. 6). While other scenarios in which the MTR complex in “Ca. Methanomixophus” could also be acquired via HGT remain possible, this condition is associated with the lowest number of MCR/MTR gene loss events during evolution (see Fig. S13a at https://figshare.com/articles/Fig_S13_Alternative_scenarios_that_might_explain_the_evolution_transition_of_Archaeoglobi_from_methanogen_/9918218). Also, the possibility that “Ca. Methanomixophus” originally inherited both the MCR and MTR complexes from the Archaeoglobi LCA cannot be excluded (see Fig. S13b). Considering the highly complex evolutionary history of the Archaea species that encode the MCR complex, recovering more MCR-encoding lineages throughout the archaean tree of life would help to make their evolutionary history and metabolic role in the hydrothermal subsurface biosphere clear.

MATERIALS AND METHODS

Sample collection and data availability. Production water was collected from the Jiangsu oil reservoir (Yangzhou, China) (18); 40 liters of the production water was collected for DNA extraction, and another 40 liters was collected for RNA extraction and was stabilized using a 10% (vol/vol) stop solution (95% ethanol, 5% TRizol [Life Technology]). All samples were kept on ice and transported to the laboratory within 4 h, and then DNA/RNA was extracted using a PowerMicrobiome RNA isolation kit (MO BIO). After sequencing on an Illumina Miseq platform was performed, metagenomic and metatranscriptomic data sets were processed as previously described (18). In brief, raw reads were quality filtered using PRINSEQ v0.20.4 (54) and were then assembled using SPAdes v3.7.0 (55), and MAGs were generated by Groomer v1.56. Newly assembled Archaeoglobi MAGs, including MAGs of “Ca. Polytropus marinifundus” (Juan de Fuca Ridge, Northeast Pacific Ocean), Archaeoglobi WYZ-LMO1 (Washburn Spring, WY, USA), and WYZ-LMO2 (Obsidian Pool, WY, USA), were retrieved from previous studies (7, 17).

Metagenome assembly and population genome binning. Metagenomes of water samples from Great Boiling Spring, NV (IMG-ID: 3300000106), and Yellowstone National Park, WY (IMG-ID: 3300058860), were downloaded from IMG/M ER and were trimmed for quality control and adapter removal using Trim_galore (http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/). The processed reads were de novo assembled using SPAdes v3.7.0 (55) with a ‘-meta’ model and with different k-mer settings (47, 79, 93, 127). The contigs generated by the different k-mer settings were evaluated using MetaQUAST v2.2 (57), and the k-mer of 127 was found to be the best (see Table S1 in the supplemental material). Scaffolds were then binned into population genomes using MetaBAT2 v0.32.4 with default settings (58).

Genome annotation. The annotation of scaffolds in population genomes was confirmed using two parallel methods. First, nucleotide sequences of genomes were submitted to theRAST server for annotation using subsystem technology (59). Second, nucleotide sequences of genomes were translated into amino acid sequences using Prodigal v2.6 (60) with default settings, and the amino acid files were submitted to BlastKOALA server (61) in the prokaryotic species database for assigning knockout (KO) numbers. Only genes with accordant annotations from both methods were included in this study. To characterize the mobile elements in the population genomes, we also searched genomes for signatures of known integrons and transposons. A local database of integrons was created from the nucleotide sequences for all integrases available in the database INTEGRALL v1.2.8414 (10,533 records in total) (62). A gene was recognized as an integron or insertion if the BLAST hit (blastn) had a minimum of 30% identity over 75% of the gene length, according to the previously published threshold (7). Amino acid files of population genomes were submitted to the iSfinder online server (updated on 4 March 2019) (63) for searching for transposons using the ‘blastp’ tool (identity > 30%, coverage > 75%, E value < 1 × 10^-5).

Construction of consensus tree. For the phylogenetic analysis of functional marker proteins (McrABG, MtrABCDHE, Sat, DsrAB, AprAB, and NarGHI), amino acid sequences of individual genes were extracted from assembled genome bins and reference genomes listed in Table S2. Sequences were aligned using MAFFT (64) with iterative refinement methods (‘G-INS-i’) and then refined (retained columns with <10% gaps), and alignments of subunits for the same enzyme were concatenated in a single alignment to get a higher level of phylogenetic resolution. To construct the genome tree, all reference genomes and assembled genome bins were pooled into PhyloPhAn v0.99, which extracts and aligns 400 conserved protein sequences from the genomes (65). The concatenated alignment file was
then extracted for phylogenomic tree building. Consensus trees were built based on three different methods similar to those described in a previous study (14). Maximum likelihood (ML) trees were reconstructed using IQ-tree v. 1.6.7 under standard conditions of model selection with 1,000 ultrafast bootstraps. Neighbor joining (NJ) trees were calculated in the PHYLIP software package (66) using the “NEIGHBOR” function based on the JTT matrix model (67), and bootstrap analysis was performed with 1,000 resamplings (PHYLIP SEQBOOT). Maximum parsimony (MP) trees were constructed in MEGA 7 (68) with 100 and 500 bootstrap replications for the genome tree and gene trees, respectively. The three trees were then combined into a consensus tree by using the extended majority rule in PHYLIP CONSENSE (66). Branch lengths of the consensus tree were inferred by using the JTT matrix model (PHYLIP PROML) (66).

Metatranscriptome analysis. Metatranscriptomes of sample W15 obtained from the Jiangsu oil reservoir were processed as previously described (18). In brief, raw reads were trimmed by quality analysis using Prinseq (with parameters identical to those used in the metagenome analysis), and the quality-controlled reads were mapped to coding DNA sequences (CDS) of the whole assembly file which has been used to generate the population genomes (18) using Bowtie2 (69) with default settings. Mapping reads are then filtered for MapQ values of ≥2 in order to remove ambiguously mapping reads (70). eXpress v1.5.1 (71) was used to calculate FPKM (fragments per kilobase per million fragments mapped).

Data availability. Amplicon sequences were deposited in the NCBI database under accession numbers MN891846 to MN891915. The genome files assembled in this study are available in the RAST server with the IDs mentioned in Table 1 and are also attached here as Text S2 in the supplemental material.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

TEXT S1, DOCX file, 0.05 MB.

TEXT S2, DOCX file, 1.3 MB.

TABLE S1, DOCX file, 0.02 MB.

TABLE S2, DOCX file, 0.02 MB.

TABLE S3, DOCX file, 0.01 MB.

TABLE S4, XLSX file, 0.02 MB.

TABLE S5, DOCX file, 0.01 MB.

TABLE S6, XLSX file, 0.02 MB.

TABLE S7, XLSX file, 0.02 MB.

TABLE S8, XLSX file, 0.04 MB.

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