



Aerobic bacterial methane synthesis

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Reports of biogenic methane (CH₄) synthesis associated with a range of organisms have steadily accumulated in the literature. This has not happened without controversy and in most cases the process is poorly understood at the gene and enzyme levels. In marine and freshwater environments, CH₄ supersaturation of oxic surface waters has been termed the “methane paradox” because biological CH₄ synthesis is viewed to be a strictly anaerobic process carried out by O₂-sensitive methanogens. Interest in this phenomenon has surged within the past decade because of the importance of understanding sources and sinks of this potent greenhouse gas. In our work on Yellowstone Lake in Yellowstone National Park, we demonstrate microbiological conversion of methylamine to CH₄ and isolate and characterize an *Acidovorax* sp. capable of this activity. Furthermore, we identify and clone a gene critical to this process (encodes pyridoxylamine phosphate-dependent aspartate aminotransferase) and demonstrate that this property can be transferred to *Escherichia coli* with this gene and will occur as a purified enzyme. This previously unrecognized process sheds light on environmental cycling of CH₄, suggesting that O₂-insensitive, ecologically relevant aerobic CH₄ synthesis is likely of widespread distribution in the environment and should be considered in CH₄ modeling efforts.

methane | aerobic | methylamine | glycine betaine | bacteria

The “methane paradox” is well documented and has drawn significant attention from a broad array of biologists, biogeochemists, biochemists, and physical scientists (see review in ref. 1). This supersaturation phenomenon is inconsistent with the extreme oxygen sensitivity of methanogens, a phylogenetically constrained group of archaea, which have long been viewed to be the sole source of biogenic CH₄. Work in marine waters previously suggested this is due to methanogens active within anoxic microhabitats in suspended organic debris (2–5), digestive tracts, or fecal pellets of zooplankton (5–7) and/or of fish (8, 9). Recently, strong evidence for microbial dealkylation of methylphosphonate (referred to here as MPn) in marine (10, 11) and freshwater environments (12, 13) has been presented and thus provides at least a partial explanation for this phenomenon.

Pelagic methane enriched zones (PMEZ, ref. 12) are well-defined CH₄ maxima occurring as a distinct region(s) of the water column in deep freshwater lakes (1, 12, 14). PMEZ offer a tractable environment in which to study the microbiology and biogeochemistry that underpins aerobic CH₄ synthesis. Our prior efforts on Yellowstone Lake using ¹³C-labeled methanogen substrates, 16S ribosomal RNA (rRNA) gene based microbial community characterization, and methyl-coenzyme M reductase (*mcrA*) gene-targeted PCRs demonstrated the absence of recognizable methanogens in PMEZ waters (12). Instead, MPn metabolism was strongly associated with distinct populations of *Pseudomonas* sp. as being important contributors to PMEZ formation (12).

Efforts presented here summarize continuing studies that now identify aerobic metabolism of methylamine (MeA) as an important contributing metabolite to biogenic CH₄. Methylotrophic methanogens anaerobically convert MeA to CH₄ by disproportionation of

MeA to carbon dioxide and CH₄ (15, 16). Methylotrophic bacteria can metabolize MeA as a carbon source (17, 18), nonmethylotrophs for nitrogen (17, 19) (20), a carbon and N source (21), or methylotrophs can use it as an energy source (22). MeA utilization is viewed to occur as an oxidation via either MeA dehydrogenase, MeA oxidase, or involving methyl group transfer to tetrahydrofolate involving the formation of formation of γ -glutamyl-methylamide (GMA) and N-methylglutamate (NMG) (20, 23). In the current study, we report the discovery of an alternate, simpler route of MeA metabolism that yields CH₄, involving a reaction catalyzed by a 5' pyridoxal-phosphate-dependent aspartate aminotransferase.

Results

Water Column Characterization. Yellowstone Lake water columns sampled in 2016 shared similar EXO Sonde features (*SI Appendix, Table S1 and Fig. S1*). The PMEZs were located just below the approximate upper limit of the thermocline, although the July PMEZ was 0.5 m deeper (11.5 versus 11 m), and CH₄ concentrations were approximately twofold greater (51 versus 26 nM) (Fig. 1A). Incubating July 29 PMEZ water samples (12 °C, initiated as aerobic) with ¹³C-labeled MPn and ¹³C-labeled substrates capable of supporting known methanogenesis pathways (acetate, formate, bicarbonate, H₂ + bicarbonate) found no strong evidence of ¹³CH₄ formation (*SI Appendix, Fig. S2A*). Samples spiked with ¹³C-MeA produced ¹³CH₄ and to a much lesser extent ¹³C-methionine as well (Fig. 1B and *SI Appendix, Fig. S2*). Lack of MPn metabolism was not anticipated and thus to relocate the MPn active region, follow-up sampling (August 5) of the PMEZ as well as 1 m above and below were spiked with

Significance

Observations summarized herein contribute to an ongoing paradigm shift in microbial ecology, documenting an emergent property of ecosystem function that further challenges the perception that biogenic methane (CH₄) production is strictly an anaerobic process. Relevant metabolites, a model bacterial isolate, gene, and enzyme are identified, and we show how this property can conceivably be broadly distributed in the biosphere and contribute to global CH₄ emissions. Scientifically, this study will enable lines of investigation that will expand our understanding of CH₄ synthesis and emission in nature and illustrates how CH₄ synthesis may actually serve as a nexus for the C and N cycles in nature.

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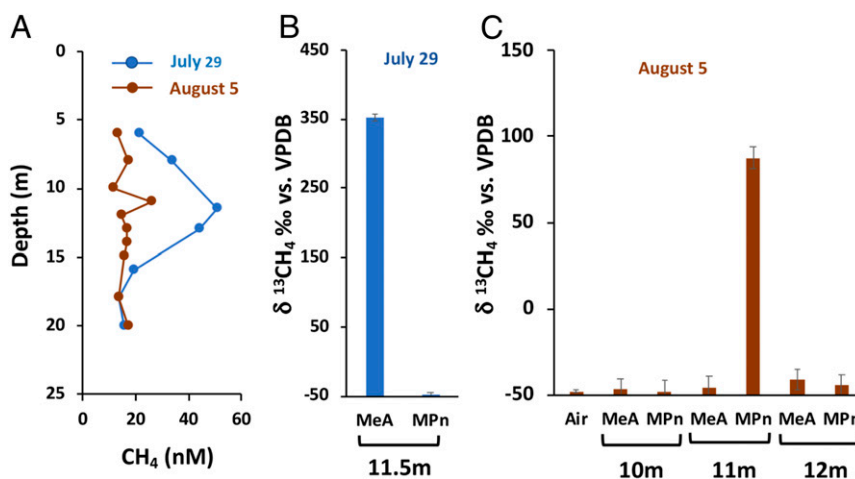


Fig. 1. Characterization of the water column methane and MPn and MeA metabolism potential of the PMEZ microbial communities. (A) Methane profiles generated from equilibrium CH_4 measurements. (B and C) ^{13}C generation from lake water samples spiked with ^{13}C -labeled methylamine (MeA) or methylphosphonate (MPn). Data are mean \pm range of $n = 2$ distinct water samples. On July 28, 2016, only water from 11.5 m was tested, whereas on August 5, 2016, water from three depths (as shown) was tested. VPDB, Vienna Pee Dee Belemnite; $\delta^{13}\text{C}$.

^{13}C -MeA, ^{13}C -methionine, and ^{13}C -MPn. On this second sampling date, ^{13}C -MPn was the primary substrate leading to ^{13}C (Fig. 1C and *SI Appendix, Fig. S2B*) and thus consistent with our prior observations (12). The data showing that MeA was the sole contributing metabolite to PMEZ formation in July was striking. MeA is widespread in the environment (24), but much remains to be learned about its environmental fate, even though it is ubiquitous and viewed as an important nitrogen source in marine environments (19, 25). Subsequent work focused on its ecological relevance in this freshwater environment.

On July 24, 2017, the water column was sampled in higher resolution (0.5-m increments). As in 2016, the PMEZ was well defined (Fig. 2A) just below the upper boundary of the thermocline (8 m) in O_2 -saturated water (Fig. 2C). Also, a second more-robust PMEZ was present at 11.5 to 12.0 m depths (roughly 3,000% of saturation) (Fig. 2A) at near O_2 saturation (98.6% of saturation, Fig. 2C). Neither PMEZ could be linked with the Chl *a* peak (14 to 16 m), and only the upper PMEZ was located within the photic zone (0 to 9 m). PCRs using degenerate universal *mcrA* primers (*SI Appendix, Table S2*) and total DNA failed to generate amplicons (*SI Appendix, Fig. S3*), indicating recognizable methanogens (26) were absent or below detection.

Targeted liquid chromatography–mass spectrometry (LC–MS) analyses of unamended PMEZ samples found MPn was below detection (1 μM in this lake water matrix; *SI Appendix, Fig. S4*). However, MeA was present ($\sim 2.8 \mu\text{M}$ at 6 m) and decreased to or below detection at depths greater than 10 m (Fig. 2B). The lowest MeA levels coincided with the deepest and largest PMEZ maxima, suggesting MeA consumption may have contributed to its formation. Results of other parallel experiments examining microbial community metabolic potentials supported this hypothesis. ^{13}C -MPn dealkylation activity indicated MPn could contribute to ^{13}C generation in water depths of 8, 11.5, and 12 m, whereas ^{13}C -MeA exhibited the potential to contribute to ^{13}C synthesis only at 12 m (Fig. 2D).

Methylated amines are indicator metabolites of glycine betaine (GB) degradation and are commonly associated with algae (27–30). Microbial conversion of GB to trimethylamine (TMA) has been documented (30, 31), which can then be converted to trimethylamine oxide (TMAO) \rightarrow dimethylamine (DMA) \rightarrow MeA (17), and DMA can be converted directly to MeA (32). In this study, GB, TMA, and MeA profiles were correlated ($R^2 = 0.64$ to 0.77, *SI Appendix, Fig. S5*), diminishing to or below the detection limits at depths that correspond to the deeper PMEZ

associated with ^{13}C -MeA \rightarrow ^{13}C activity (Fig. 2B and D). This suggests that the source, consumption, and metabolism of the three metabolites could be coordinated.

Microbial Community and Underlying Biology. Analysis of Illumina 16S rRNA gene sequence libraries was used to determine if specific taxa exhibited increased abundance at the 12-m depth PMEZ and hence might indicate which organism(s) could be involved in ^{13}C -MeA \rightarrow ^{13}C activity. Taxa classified as belonging to the order *Burkholderiales* (annotated as Acid A-1 and Acid A-2 in the Freshwater Database) exhibited a sharp peak (13% abundance, Fig. 2A) coinciding specifically with the MeA metabolic potential at 12 m (Fig. 2D). However, in agreement with the apparent absence of amplifiable *mcrA* (*SI Appendix, Table S2*), recognizable methanogen signatures were absent in all libraries throughout the water column.

To more closely examine the biology underpinning the MeA \rightarrow CH_4 activity, microorganisms from PMEZ samples were isolated on SAR-MeA agar containing MeA as the sole nitrogen source (*Methods*) and then screened in liquid SAR-MeA for CH_4 generation. We isolated several bacteria capable of converting MeA to CH_4 . Genus-level classification based on 16S rRNA gene sequence includes the following: *Acidovorax* sp. (MK896843.1), *Pseudomonas* sp. (MK896839.1), *Caulobacter* sp. (MK896844.1), *Mesorhizobium* sp. (MK896845.1), and *Dietzia* sp. (MK896847.1). For further characterization, we selected one of the *Acidovorax* sp. isolates obtained from the 12-m depth because this genus is a member of the *Burkholderiales* order that was significantly abundant at the depth corresponding to the MeA metabolic potential (Fig. 2A and D) and because it was found to be genetically tractable. Batch cultures in liquid SAR-MeA in sealed serum bottles (maintained aerobic with sterile air injections at each gas sampling) demonstrated robust growth (*SI Appendix, Fig. S6A*) and CH_4 generation from MeA (Fig. 3A). Gas chromatography–mass spectrometry (GC–MS) analysis of headspace samples of the *Acidovorax* sp. cultured with ^{13}C -MeA confirmed ^{13}C generation (*SI Appendix, Fig. S7*), demonstrating transfer of the MeA carbon to CH_4 . GB was also investigated in the context of the above hypothesized catabolic pathway and also found to support growth (*SI Appendix, Fig. S6A*) and CH_4 generation (Fig. 3A).

Gene Identification and Mutational Analysis. Transposon Tn5B22 mutagenesis of the *Acidovorax* sp. isolate yielded two mutants (among $\sim 8,000$ transconjugates) capable of growth with ammonium

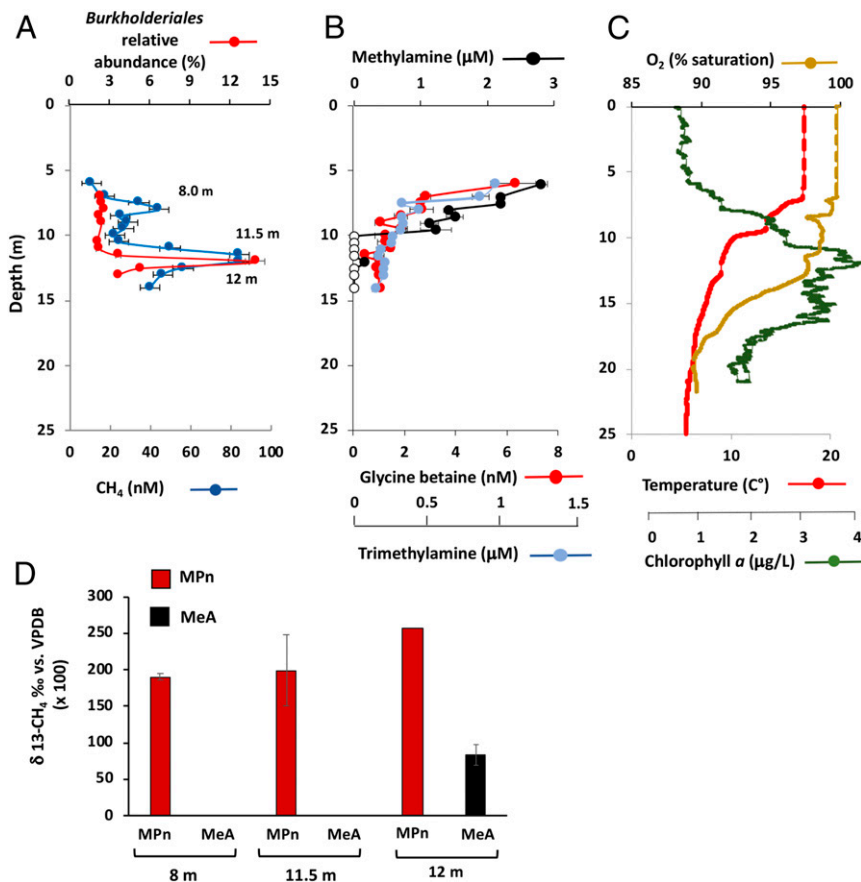


Fig. 2. PMEZ characterization in 2017 for methane and relevant catabolic precursors. (A) Water column methane profile showing two concentration maxima at 8.0 and 11.5 to 12.0 m overlaid with relative abundance (%) of *Burkholderiales*. (B) Metabolite analysis showing change in concentrations of glycine betaine, trimethylamine, and methylamine. Open symbols indicate methylamine concentration below the detection limit. (C) Water column properties illustrating O_2 saturation (or nearly so) at both methane peak depths. (D) Relative metabolic potentials of the microbial communities for converting ^{13}C -MPn or ^{13}C -MeA to ^{13}C - CH_4 . Data in D denotes the mean \pm range of two replicates except for MPn at 12 m, at which $n = 1$ due to broken serum bottle. VPDB, Vienna Pee Dee Belemnite; $\delta^{13}C$, ratio of stable isotopes ^{13}C : ^{12}C in parts per thousand (per mil, ‰).

chloride but not MeA nor GB as sole N source (*SI Appendix, Fig. S6B*) and were likewise defective for generating CH_4 (Fig. 3B). Cultures initiated at cell densities equivalent to mid log phase under

these growth conditions (starting optical density = 0.15; *SI Appendix, Fig. S6*, compare D and E) also failed to generate CH_4 , demonstrating lack of CH_4 synthesis in the mutants is due to defective

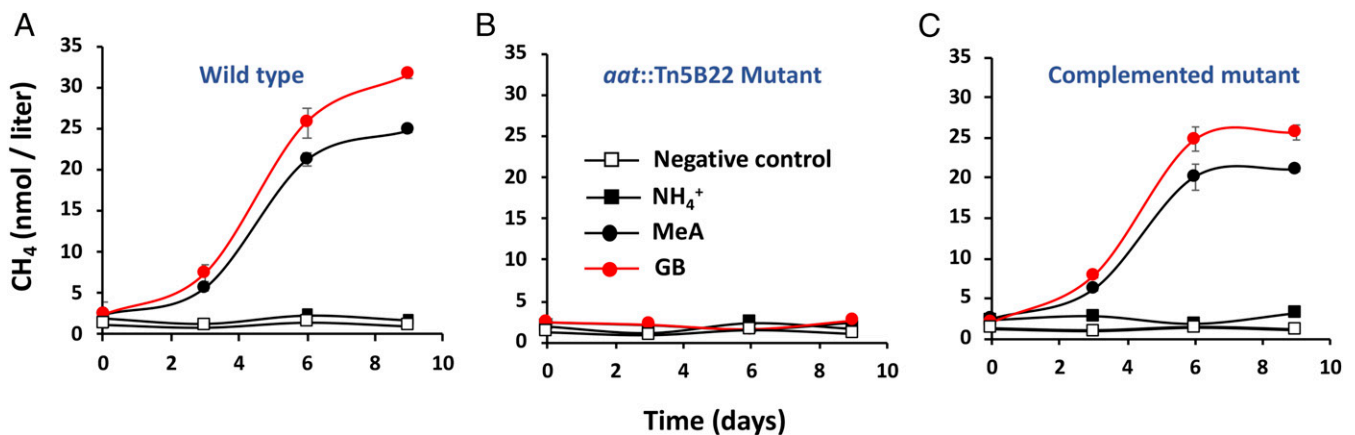


Fig. 3. Conversion of glycine betaine and MeA to CH_4 by pure culture *Acidovorax* sp. isolates. CH_4 generation profiles are shown for the following: (A) wild-type *Acidovorax* sp. isolate; (B) the *aat::Tn5B22* mutant 3-29; and (C) 3-29 mutant carrying the pCPP30::*aat* that complements the mutant back to wild type for growth on glycine betaine and MeA and for CH_4 synthesis. All strains were cultured in SAR media containing NH_4^+ , MeA, or glycine betaine as the sole nitrogen source (treatment symbols for all three panels are shown in B). Data points and error bars (where visible) are the average \pm SD of three replicates.

MeA and/or GB metabolism, not lack of biomass. The two Tn5B22 insertion sites differed, but the same gene was affected in both mutants (*SI Appendix, Fig. S8*), providing strong evidence that the mutated gene is essential to MeA/GB metabolism and CH₄ synthesis in this organism. Basic local alignment search tool (BLAST)x analysis matched the gene to the family of pyridoxal 5' phosphate-dependent aspartate aminotransferases (referred to herein as *plp-aat*). The specific *plp-aat* coding sequence without adjacent genomic DNA (GenBank accession no. MK170382) was PCR cloned from the *Acidovorax* isolate DNA into the broad host range plasmid pCPP30. Conjugation of the recombinant plasmid to the mutants reversed their negative growth and CH₄ synthesis phenotypes (Fig. 3C and *SI Appendix, Fig. S6C*), confirming the importance of this gene.

Introducing *plp-aat* to *Escherichia coli* strain BL21(DE3) allowed this enteric bacterium to grow in M9 broth containing MeA as the N source (Fig. 4A) and to synthesize CH₄ (Fig. 4B). However, growth in M9-MeA media required BL21 (pET28::*aat*) to first be grown in lysogeny broth (LB) broth with antibiotic selection; that is, washed cells failed to grow in M9-MeA. Consequently, subsamples of stationary phase cells were transferred directly (~200-fold dilution) to the M9-MeA media (with isopropyl β-d-1-thiogalactopyranoside [IPTG] to induce *aat* transcription). The identity of the putative metabolite(s) apparently necessary for the recombinant *E. coli* to grow in M9-MeA is unknown at present, but it was not sufficient to allow the BL21 (pET28a+) negative control to grow in M9-MeA. Endpoint oxygen levels for the *E. coli* experiments were ~43% of saturation at 37 °C (*SI Appendix, Fig. S9*), illustrating the cultures had not gone anaerobic. For both *Acidovorax* (Fig. 3 and *SI Appendix, Fig. S6A*) and *E. coli* (Fig. 4), CH₄ synthesis was linked to growth on MeA, implying N acquisition and CH₄ release are somehow metabolically coupled.

PLP-Aat enzyme has been used as a model enzyme for understanding PLP-linked catalysis, and from these efforts, an essential catalytic lysine residue has been identified to be invariantly conserved (33, 34). Prior work with the *E. coli* PLP-Aat enzyme has shown that mutation of this conserved lysine to alanine results in an enzyme with only 0.5% native aspartate aminotransferase activity (34). In an analogous approach, we changed the catalytic lysine (K237) to alanine in the cloned *plp-aat* and repeated the bioassays. Growth and CH₄ synthesis by BL21 carrying the K237A mutant clone was severely constrained in MeA media (Fig. 4).

Purified Enzyme Generates CH₄ from MeA. To more directly test the PLP-Aat protein for CH₄ production, it was overexpressed and purified as a His-tagged protein (*SI Appendix, Fig. S10*). For assessing PLP-Aat activity, enzyme reactions were conducted in *E. coli* cytoplasm extract that was prefiltered through a 3 kDa molecular weight (MW) cutoff filter to remove proteins but retain all low molecular weight species (*Methods*) so as to include any other metabolite(s) that are unknown at present but which might be required for *E. coli* to host this activity. When purified PLP-Aat was added to the cytoplasmic extract along with α-ketoglutarate and PLP, CH₄ was produced (Fig. 5). Control reactions using heat inactivated enzyme did not produce CH₄. Likewise, negative were reactions where α-ketoglutarate or MeA were individually omitted. Methane was produced at a lower level in reactions where PLP was omitted. We attribute this to apo-form enzyme that purified with holo-form enzyme. This could be due to overproduction of the enzyme in the pET system and is consistent with the increased activity upon addition of PLP, which would generate more holo-form enzyme (PLP-Aat).

Discussion

From experiments including analyses of microbial community composition and metabolic potential, targeted metabolomics, pure culture physiologic and genetic characterizations, and defined genetic transfer of this property, we put forth data that contributes to an ongoing paradigm shift for environmental CH₄ synthesis that is distinct from classical methanogenesis or other presently known mechanisms of CH₄ synthesis. We found MeA occurs at substantial concentrations in the pristine Yellowstone Lake environment in conjunction with GB and TMA (Fig. 2B and *SI Appendix, Fig. S5*), which are catabolic precursors of MeA. The extant microbial community exhibited the capacity to convert MeA to CH₄ (Fig. 2D), and depending on as of yet undefined environmental conditions, MeA is an important contributor to PMEZ formation (Fig. 1, Fig. 2D) or indeed the only contributor (Fig. 1A and B). The ability to convert MeA to CH₄ is heritable by a single gene (Figs. 3C and 4) and importantly, MeA→CH₄ conversion will occur under aerobic conditions (Figs. 3 and 4 and *SI Appendix, Fig. S9*). In sum, this study illustrates all the essential components of a relatively simple and ecologically relevant CH₄ synthesis pathway; that is, aerobic bacterial CH₄ synthesis.

Some have argued that lateral transport of CH₄ derived from near shore-based methanogenesis to be the basis for metalimnetic CH₄ (35–37). However, evidence supporting in situ CH₄ synthesis in oxic lake waters is now clear and overwhelming (e.g., refs. 1, 12

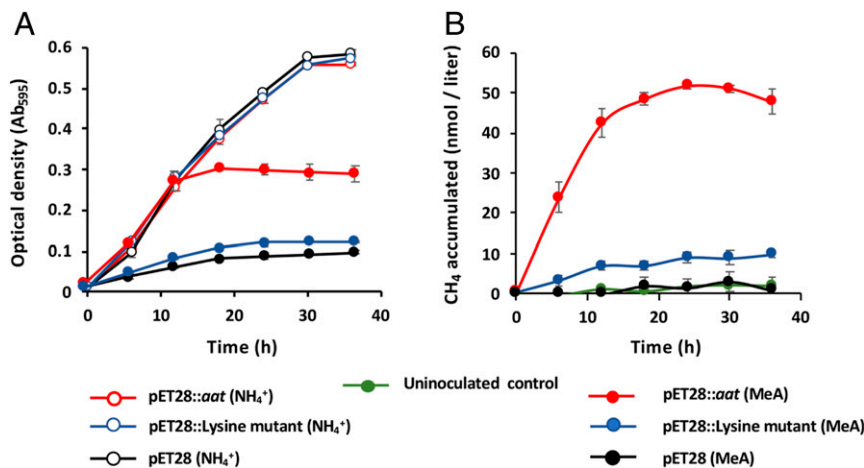


Fig. 4. PLP-Aat enables *E. coli* to grow with MeA as a nitrogen source and synthesize CH₄. (A) Growth profiles of *E. coli* BL21 with MeA or NH₄⁺ and carrying the empty plasmid (control), cloned *aat* (pET28a::*aat*), or mutant *aat* wherein the catalytic lysine (K237) was replaced by alanine. (B) CH₄ generation by BL21 with the same constructs. All data and error bars (where visible) are the mean ± SD of three replicate cultures.

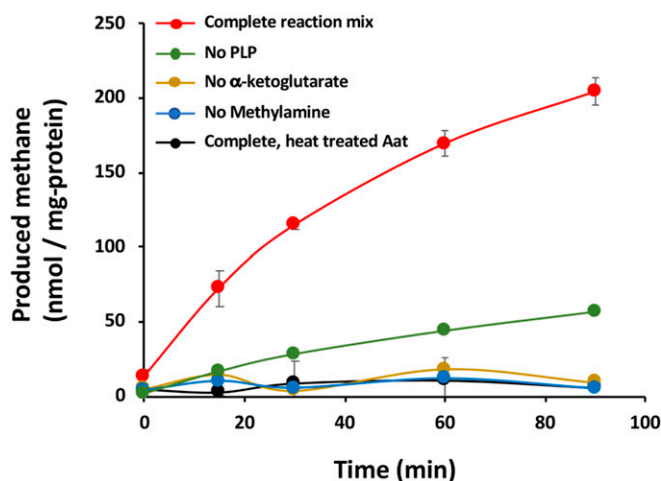


Fig. 5. CH_4 synthesis by purified Aat. CH_4 synthesis by purified Aat in *E. coli* cytoplasmic extract. Complete reactions contained purified enzyme (typically $\sim 3 \mu\text{g}$), MeA (4 mM), α -ketoglutarate (4 mM), and 5'pyridoxal phosphate (1 mM). Other reactions shown were conducted wherein a single substrate was omitted in order to illustrate the requirement of each for the generation of methane. All data are the mean \pm SD of three replicate reactions.

to 14, 38, and 39). We suggest that defined CH_4 maxima (PMEZs) that coincide with relevant microbes (ref. 12 and Figs. 2 and 3), substrates (ref. 12 and Fig. 2B), and functional gene occurrence (12) is not coincidental nor explainable by lateral transport. Furthermore, recent work directly comparing in situ oxic CH_4 production versus lateral transport concluded that oxic biogenic CH_4 accounts for the majority of CH_4 emissions for freshwater lakes with surface areas $> 1 \text{ km}^2$ (40). As such and given estimates that freshwater CH_4 release accounts for roughly 16% of total annual global emissions (41), aerobic CH_4 synthesis from GB and or MeA may contribute substantially to biosphere CH_4 emissions.

While substrates supporting CH_4 synthesis likely derive from photosynthetic primary producers in aqueous environments (42, 43), the Yellowstone Lake PMEZ is consistently (across four sampling seasons) not associated with a Chl *a* peak (e.g., Fig. 2A and SI Appendix, Fig. S1). Subsurface chlorophyll maxima result from phytoplankton sinking/migration behaviors associated with nutrient depletion or from photoadaptation at deeper locations (44). Metabolite exchange between sinking phytoplankton and bacteria (45–47) may be an active (potentially mutualistic) process or derived from algal cell lysis caused by bacterial algaeacides or virus lytic events (48–50). Increased water density at colder temperature transitions in the water column could result in transient accumulation of substrates (i.e., live functioning or lysed algae, particulate organic matter) (51) at specific depths. Enhanced residence time would facilitate CH_4 synthesis and accumulation, resulting in PMEZ formation. CH_4 concentration at these depths does not necessarily correlate with synthesis rate and may be more easily explained as synthesis exceeding consumption. The absence of dissolved MPn does not necessarily mean that alkylphosphonates are absent (detection limit $1 \mu\text{M}$, SI Appendix, Fig. S4). Our assay methodology would not detect methylphosphonate (or any alkylphosphonates) covalently bound to organic matter in marine water (10). Indeed, the microbiome metabolic capacity to convert MPn to CH_4 was present in both PMEZs and consistently at the more shallow PMEZ. Presumably, to some extent the occurrence and separation of MPn and MeA metabolic potentials in the water column is tied to bioavailability of relevant substrates. Methionine conversion to CH_4 was limited relative to MPn and MeA but nevertheless detectable. Prior studies have documented methionine conversion to CH_4 , although under anaerobic conditions (52–54).

We predict compounds such as GB as well as choline (55–57) play an important role in generating methylated amines and thus are likely to contribute to this aerobic CH_4 synthesis pathway, particularly in marine environments. GB is a ubiquitous osmolyte in bacteria, archaea, and eukaryotes (23, 58) and a precursor to the equally ubiquitous MeA, which is also recognized as an important N source (19, 23, 59) occurring at concentration ranges from nM to μM (60–64). Methylated amines are indicator metabolites of GB degradation (28), are common in algae (6, 32) or phototrophic mats (28, 29), and consistent with the documented biochemistry and genetics underlying GB conversion to TMA and TMA conversion to MeA in various organisms (17) (30, 31) (32). MeA metabolism has likewise been examined in some detail (23, 25), although this prior characterization work did not include synthesis of CH_4 as an end product.

Ecological-based evidence in support of an aerobic $\text{GB} \rightarrow \text{CH}_4$ pathway includes research which infers CH_4 production associated with methylamines derived from zooplankton grazing of phytoplankton (6) and recent studies showing TMA enhances CH_4 synthesis in N-depleted incubations of Lake Stechlin oxic PMEZ water samples (65). Furthermore, the current study provides direct genetic evidence as illustrated by the inability of *Acidovorax aat::Tn5B22* mutants to synthesize CH_4 from GB nor use GB as an N source (Fig. 3B and SI Appendix, Fig. S6 B and E). By contrast, the mutants were reverted back to wild-type status when provided with the *plp-aat* gene (Fig. 3C). This indicates that, at least for this organism, acquiring N from GB must include the biochemical step catalyzed by the enzyme encoded by *plp-aat* described herein.

Several studies speak to the general importance of GB and MeA beyond simply their ubiquity and abundance in marine environments. Potential fates include assimilation as a carbon source, oxidized for energy, or used as a nitrogen source (19, 22, 25, 66–68). These examples represent metabolisms that would compete with a $\text{MeA} \rightarrow \text{CH}_4$ pathway, but it is still reasonable to consider its potential for CH_4 synthesis. At the moment, information on marine MeA dynamics is limited, but MeA assimilation (0.005 to $54 \text{ nmol L}^{-1} \text{ day}^{-1}$), oxidation (0.09 to $3.43 \text{ nmol L}^{-1} \text{ day}^{-1}$) and turnover rates (40 to $57 \text{ nmol L}^{-1} \text{ day}^{-1}$) have been reported (60, 67), facilitating at least a preliminary estimate. Conservatively assuming a PLP-Aat catalyzed MeA turnover rate of $1 \text{ nmol L}^{-1} \text{ day}^{-1}$ in the top 100 m of the global oceans is associated with CH_4 release, this would account for 11.6% of annual marine CH_4 emission [$2.1 \text{ Mt CH}_4 \text{ yr}^{-1}$ (69); Methods].

This mode of CH_4 synthesis could be highly relevant to any situation where GB and or MeA occur, including environments where CH_4 synthesis has traditionally been interpreted to derive from methanogens. There is no obvious reason why this PLP-Aat reaction would be constrained by redox conditions per se because aminotransferase reactions are important for maintaining cellular amino acid equilibria in all self-replicating organisms. Indeed, numerous aerobic or anaerobic organisms may be involved with varying efficiency, and at present, we are unable to offer any specific criteria for excluding hundreds of more-remote *aat* homologs in metagenome libraries that may encode the same function, perhaps with greater efficiency. However, we draw attention to the fact that the PLP-Aat encoded in the genome of *E. coli* strain BL21(DE3) (GenBank genome accession no. AM946981, used herein for bioassays) shares only 21.6% identity with the *Acidovorax* enzyme and did not allow *E. coli* to grow with MeA nor make CH_4 from MeA. Thus, apparently not all proteins annotated as PLP-dependent Aat enzymes are equal in regard to catalyzing the $\text{MeA} \rightarrow \text{CH}_4$ reaction. Characterizing homologs capable of this reaction will enhance our understanding of the universality of this process within and across environments. A cursory search identified thousands of homologs with a high degree of homology with the *Acidovorax* PLP-Aat, suggesting the possibility that $\text{MeA} \rightarrow \text{CH}_4$ activity occurs in other aqueous and terrestrial environments as well. Examples of highly homologous Aat proteins occurring in a

variety of aquatic environments are provided in *SI Appendix, Fig. S11*.

The PLP-Aat catalytic mechanism resulting in CH₄ release is unknown at present. In particular, the source of reductant represents a mechanistic enigma that further contributes to the methane paradox (70). Recreating the reaction milieu in *E. coli* cytoplasmic extract amended with MeA, α -ketoglutarate, PLP, and live purified PLP-Aat enzyme resulted in methane generation (Fig. 5). Methane was not generated when enzyme was heat inactivated before addition. Prefiltering the extract using a 3 kDa cutoff eliminates a direct contribution of other enzyme(s) but would not remove other metabolite(s), which may contribute or may be associated with the requirement for *E. coli* to be first grown in a complex medium. The functional importance of the catalytic lysine that is conserved across all PLP-Aat enzymes appears to be involved for the MeA \rightarrow CH₄ reaction and thus implies some basic similarities; that is, the Schiff base formation between the Aat catalytic lysine and the aldehyde carbon of the coenzyme pyridoxal 5'-phosphate (33, 34). However, beyond this initial step it is also not possible at this juncture to predict reaction mechanism, or efficiency, or any other metabolite/gene product apparently involved in the functioning cell, though it is relevant to point out that CH₄ synthesis in both *Acidovorax* (Fig. 3) and *E. coli* (Fig. 4) appears connected to growth, implying a link between CH₄ release and N acquisition to support growth. And furthermore, for at least organisms such as the *Acidovorax* sp. isolate characterized herein, growth with MeA was as robust as ammonium (*SI Appendix, Fig. S6*), arguing that MeA use as a nitrogen source does not represent a spurious metabolism. Indeed, *aat::Tn5B22* mutants failed to grow with MeA, which argues that alternate MeA metabolic pathways (23) are missing in this organism, which is also consistent with the absence of genes annotated as encoding N-methylglutamate synthase, γ -glutamylmethylamide synthase, and N-methylglutamate dehydrogenase in the draft genome of this organism.

Establishing the essential components of a GB \rightarrow MeA \rightarrow CH₄ pathway in a pure culture of an ecologically relevant microorganism and linking this to in situ metabolite analyses and community metabolic potentials reveals a route of CH₄ synthesis occurring under fully oxygenated conditions. This also elucidates a nexus for C and N cycling in nature. Based on the characterization of the *Acidovorax* sp. isolate detailed herein, we view this organism as a robust, though almost certainly not unique, participant in PMEZ formation and useful as a first-generation model for illustrating gene–enzyme–organism–ecosystem linkages in this process.

Methods

Study Site, Sampling, and Initial Sample Analyses. The Yellowstone Lake sampling location and YSI EXO1 multiparameter Sonde water column characterizations were as previously described (12). For 2016 samplings, PMEZ localizations were determined using a dissolving gas equilibration system also as described (12). In 2017, the more extensive water column sampling did not allow for the use of the more time-consuming flow-through gas equilibration device for PMEZ identification. Instead, the thermocline was identified and characterized using the EXO Sonde, which then allowed us to approximate the depth of the anticipated PMEZ (i.e., just below the upper limit of the thermocline) (12). Extensive sampling was then conducted above and below this depth. On the day of sampling in 2017, lake conditions were exceptionally placid, allowing for sampling in 0.5-m increments.

For in situ CH₄ analysis, duplicate lake water samples for each depth were collected in 250 mL serum bottles, filling from the bottom and overflowing with at least one volume to expel bubbles. Vials were immediately sealed with gray chlorobutyl rubber stoppers and secured with aluminum crimps. Upon reaching shore, samples were killed by injecting 200 μ L saturated HgCl₂ solution and stored on ice for transport to the laboratory where they were stored at 4 °C. CH₄ analysis was by GC of an introduced headspace (using ultra-high purity N₂) on a Hewlett-Packard HP5890A (for 2016 work) or a Varian CP-3800 (for 2017 work) gas chromatograph, both with flame ionization detection. The original CH₄ concentration in solution was calculated using Henry's law and solubility equations (71). All other water samples were stored on ice in the dark for transport to the laboratory and then stored in a cold room (5 °C).

Upon identifying the PMEZs from the GC analysis (~24 h post-lake acquisition), untreated cold room stored samples were then used to initiate enrichments with ¹³C-labeled substrates to identify and qualitatively assess potential methanogenesis substrates yielding ¹³CH₄. For the 2016 samples, select water column samples were aseptically spiked to a final substrate concentration of 1 mM with filter sterilized stock solutions (25 mM, 99 atom%, Sigma-Aldrich) of ¹³C-labeled formate, acetate, NaHCO₃, NaHCO₃ + H₂, MPn, MeA, or methionine. PMEZ samples taken in 2016 were incubated in the dark at 12 °C to mimic in situ conditions. ¹³C enrichments of samples taken in 2017 focused only on ¹³C-MPn and ¹³C-MeA. Also, by that time we had found all functionally relevant lake isolates to have mesophilic temperature optima and thus incubations were conducted at room temperature, which was more optimal for assessing the potential of lake microbial community samples to convert these substrates to ¹³CH₄. The isotopic signature ($\delta^{13}\text{C}$) of the CH₄ of the headspace was measured using a modification of the protocols described by Wang et al. (12) on a Picarro G2201-*i* cavity ring-down spectrometer (CRDS) equipped with a Small Sample Introduction Module 2 (SSIM2). Each vial was first pressurized with 20 mL of <0.2 μ m filtered ambient air ($\delta^{13}\text{C}$ = ~-48‰ versus Vienna Pee Dee Belemnite) and then 20 mL was removed via gastight syringe and injected into the SSIM2. Factory calibration was used for the CRDS, and a tank of compressed ambient air was used as a reference material between injections to verify lack of instrument drift. For samples taken in 2017, the isotopic signature ($\delta^{13}\text{C}$) of the CH₄ of the headspace was determined using GC-MS.

GC-MS Analysis. For CH₄ isotope analysis, 800 μ L headspace gas was injected directly into an Agilent 7890 GC-MS equipped with Carboxen-1010 porous-layer open-tubular (PLOT) capillary GC column in splitless mode. The injection temperature was 200 °C, the ion source was set to 230 °C, helium was the carrier gas, and the column flow rate was 1 mL/min. The oven was held at 35 °C for 7.5 min and then heated at 25 °C/min to 250 °C and held at that temperature for 6.5 min. The mass spectrometer was operated in scan/selected ion monitoring (SIM) mode; scan range (10 to 100 *m/z*) for SIM *m/z* 14, 15, 16, 17 were used. The ¹³CH₄ standard was purchased from Sigma-Aldrich, and for ¹²CH₄, a standard high purity methane tank was purchased from American Welding and Gas.

For analysis of *E. coli* culture headspace oxygen, 250 μ L headspace gas was injected directly into an Agilent 7890 GC-MS equipped with a Carboxen-1010 PLOT capillary GC column in splitless mode. The injector temperature was 200 °C, with the ion source at 230 °C and a helium flow rate of 1 mL/min. An isocratic GC program at 65 °C with the mass spectrometer in SIM mode for *m/z* 32 was used.

Detection of GB, TMA, and MeA. Methylamine hydrochloride, trimethylamine hydrochloride, and glycine betaine hydrochloride standards were purchased from Sigma-Aldrich. For detection of MeA in lake waters, dansyl chloride (Dns-Cl, 1-dimethylaminonaphthalene-5-sulfonyl chloride) was used to label the compounds of interest prior to analysis by LC-MS. The dansylation procedure was performed as described by Guo et al. (72). Briefly, a Hamilton gastight syringe was used to draw 50 μ L water from the sample serum bottle, which was then placed into a 250 μ L polypropylene analysis vial and pH adjusted to ~9.5 with 2 μ L 160 mM sodium hydroxide. Dns-Cl prepared in acetonitrile (20 mg/mL) was added to the sample in a volume of 46 μ L. Samples were then incubated for 30 min at room temperature. After the incubation period, pH was adjusted to ~4 with 2 μ L 10% formic acid. At this point, the sample was ready for analysis. Chromatography experiments were done on an Agilent 6538 quadrupole time-of-flight (Q-TOF) mass spectrometer, positive mode, equipped with a reversed-phase Agilent Zorbax Eclipse Plus C18 column (2.1 \times 150 mm). Solvent A was 0.1% formic acid in high-performance liquid chromatography (HPLC) water, and solvent B was 0.1% formic acid in acetonitrile. The 15 min binary gradient elution profile was as follows: *t*₁) 1 min, 0% B; *t*₂) 11 min, 55% B; *t*₃) 14 min, 100% B; and *t*₄) 15 min, 0% B. The wavelength was 320 nm, the flow rate was 600 μ L/min, and the sample injection volume was 10 μ L. Limit of detection was determined to be 20 nM.

For detection of trimethylamine, ethyl bromoacetate was used to label the compound of interest prior to analysis by LC-MS. The labeling procedure was performed as described by Johnson (73). A Hamilton gastight syringe was used to draw 90 μ L water from the sample serum bottle and transferred to a 250 μ L polypropylene analysis vial and then 10 μ L (20mg/mL acetonitrile) ethyl bromoacetate was added. Samples were then incubated for 30 min at room temperature. At this point, the sample was ready for analysis. Chromatography was done on an Agilent 6538 Q-TOF mass spectrometer, positive mode, equipped with a normal-phase Waters ACQUITY BEH HILIC 1.7 μ m column (2.1 \times 100 mm). Solvent A was 0.1% formic acid in HPLC water, and solvent B was 0.1% formic acid in acetonitrile. The 4.7 min binary gradient elution

profile was as follows: t_1) 0 min, 90% B; t_2) 2 min, 70% B; t_3) 3.6 min, 60% B; and t_4) 4.7 min, 90% B. The flow rate was 400 $\mu\text{L}/\text{min}$, and the sample injection volume was 5 μL . Limit of detection was determined to be 50 nM.

For detection of glycine betaine, a Hamilton gastight syringe was used to transfer 90 μL water from the sample serum bottle to a 250 μL polypropylene analysis vial. Methanol was added to give a 10% concentration, and the sample was then vortexed for 30 s. At this point, the sample was ready for analysis. Chromatography was done on an Agilent 6538 Q-TOF mass spectrometry, positive mode, equipped with a normal-phase Waters ACQUITY BEH HILIC 1.7 μm column (2.1 \times 100 mm). Solvent A was 0.1% formic acid in HPLC-grade water, and solvent B was 0.1% formic acid in acetonitrile. The 4.7 min binary gradient elution profile was as follows: t_1) 0 min, 90% B; t_2) 2 min, 70% B; t_3) 3.6 min, 60% B; and t_4) 4.7 min, 90% B. The flow rate was 400 $\mu\text{L}/\text{min}$, and the sample injection volume was 10 μL . Limit of detection was determined to be 1 nM.

Isolate Enrichment, Cultivation, and Characterization. All bacteria used in this study are listed in *SI Appendix, Table S2*. To isolate *Acidovorax* from the lake water, we used the same SAR media described previously as SAR-MPn (12), except modified such that 10 mM glucose replaced pyruvate, 1 mM MeA or GB was provided as the sole N source instead of NH_4SO_4 , and inorganic phosphate was used instead of MPn as a P source (agar media referred to herein as SAR-MeA). Sample aliquots from the positive $^{13}\text{C}\text{-MeA} \rightarrow ^{13}\text{CH}_4$ enrichments were plated directly onto SAR-MeA solidified with Agar Noble. Following a 2-wk incubation (room temperature), visually unique colonies were identified based on morphological differences and selected for several rounds of subculture on SAR-MeA agar to obtain pure cultures.

DNA from each culture was extracted using Wizard Genomic DNA Purification Kits (Promega) as per manufacturer's instructions. The 16S rRNA genes were PCR cloned using 27F and 1492R PCR primers (*SI Appendix, Table S2*). Amplicons were individually cloned into pCR2.1 (Invitrogen); clone-bearing plasmids were purified via QIAprep Spin Miniprep Kit (Qiagen) and then sequenced by the Brigham Young University Central DNA sequencing facility. Resulting sequences were compared with GenBank sequences to identify closely related cultured organisms via BLASTn.

For culture characterizations, *Acidovorax* was grown in liquid SAR-MeA media containing glucose as the sole carbon source (4 g/L). Growth was tracked based on culture optical absorbance (Ab_{595}) using a Molecular Dynamics microtiter plate reader. MeA \rightarrow CH_4 synthesis experiments were conducted in 70 mL sealed serum bottles in order to quantify CH_4 synthesis, which was measured via GC as described in *Study Site, Sampling, and Initial Sample Analyses*.

DNA Isolation, PCR, and Sequencing. DNA was extracted from all water column depths for use as PCR templates to determine whether the methanogen indicator gene, *mcrA*, was detectable. In each case, biomass from triplicate 1 L samples were separately collected on Sterivex-GV 0.22 μm filter cartridges (Millipore Sigma) and then DNA extracted using the PowerWater DNA isolation kit (MO BIO Laboratories, Inc.) following the manufacturer's instructions.

PCR primers for near full-length amplification of the *Acidovorax* pure culture 16S rRNA gene used the primers describe by Lane (74), and for the *mcrA* gene, we used the universal primers described by Luton et al. (75) (*SI Appendix, Table S2*). For community 16S rRNA gene sequencing, DNA extracts were quantified using a SpectraMax Plus Microplate Reader (Molecular Devices) and PCR tested prior to submission for Illumina sequencing at the Institute for Genomics and Systems Biology Next Generation Sequencing Core at Argonne National Laboratory. Very briefly, PCR amplicons were generated using barcoded primers 515F and 806R targeting the V4 region of the 16S rRNA gene in the domains Bacteria and Archaea (76) and then sequenced using the Illumina MiSeq sequencing platform. Illumina sequence libraries were processed using the mothur software package [version 1.39, (77)]. Low-quality sequences were removed using minimum and maximum lengths of 100 and 500, respectively, and no ambiguous bases or mismatches were allowed in the primer sequence. Chimera sequences were removed using UCHIME (78), and the Silva v123 reference database was used to remove uninformative data (79). For each depth, the libraries were rarified to 18,800 quality and trimmed reads, and singletons were removed prior to calculating relative abundances. Operational taxonomic unit classification was based on 97% identity, and classification was based on using the "FreshTrain" database available at <https://github.com/McMahonLab/TaxAss/tree/master/FreshTrain-files> (80). Repeated resampling ($n = 10$) of the libraries generated the same relative abundances for the taxa described. Sequencing data can be found in GenBank as Bioproject ID PRJNA598368, Biosample accession no. SAMN13704654.

Transposon Mutagenesis, *pIp-aat* Cloning, and Complementation. Methods for transposon mutagenesis, genome walking, and cloning were as we previously described (81). Briefly, *E. coli* S17-1 was used to conjugate Tn5-B22 (82) to *Acidovorax* sp. Gent^R transconjugates were then screened for loss of growth on SAR-MeA agar, yielding two unique mutants. Cloning of the Tn5B22 and adjacent DNA, as well as additional primer walking, used the APAGene genome walking kit (Bio S&T Inc.) to obtain the full gene sequence. The specific *aat* sequence was then PCR cloned into pCR2.1 and then subcloned to the broad host range plasmid pCPP30, which was then transformed into *E. coli* S17-1 for conjugal transfer to each mutant for complementation experiments.

Catalytic Lysine Amino Acid Substitution Mutation. The selected genes were refactored to match the codon usage of *E. coli* using an empirically derived codon usage table to remove NdeI and HindIII restriction sites and to overcome DNA synthesis constraints using Build-Optimization Software Tools (BOOST) (83). Synthetic DNA (Twist Biosciences) was inserted into the NcoI/XhoI site of pET28a+ by Gibson assembly. The *E. coli* Top10 transformants were plated on LB agar plates supplemented with kanamycin (50 $\mu\text{g}/\text{mL}$). All plasmids were sequence verified using the PacBio Sequel sequencing platform (Pacific Biosciences). Positive clones were DNA prepped and transformed into BL21 (DE3) cells (Novagen).

Enzyme Purification. *E. coli* strain BSL21 carrying pET28a⁺::*aat* was used as the expression system for the His-tagged Aat. Briefly, cells were grown in LB media under kanamycin selection at 37 $^{\circ}\text{C}$ to an OD_{600} of 0.2 and then Aat protein production were induced by adding IPTG to a final concentration of 0.1 mM. Cells were incubated at 25 $^{\circ}\text{C}$ for 8 h, then spun down (3,000 rcf, 20 min, 4 $^{\circ}\text{C}$). The resulting cell pellet was stored at -80°C until purification. Cells were lysed using sonication in a lysis buffer of phosphate buffer (0.1 M sodium phosphate, pH 7.4), 10 mM imidazole, and cOmplete Mini EDTA-free Protease Inhibitor mixture (cOmplete, Sigma-Aldrich). Protein was purified using a step-wise imidazole gradient on a HisTrapFF affinity column (GE Healthcare). Following elution, the Aat protein fraction was dialyzed into phosphate buffer overnight using 10 k molecular weight cut off (MWCO) SnakeSkin pleated dialysis tubing (ThermoScientific). The Aat fraction was then concentrated using a Nanosep 30 k MWCO centrifugal device (PALL Life Sciences), and glycerol was added to a final concentration of 10%. Protein was stored at -80°C until used in the enzymatic assays. Intact protein molecular weight determination was conducted using C4 reverse-phase LC-MS on Micro-TOF (Bruker Daltonics) coupled to a 1290 ultra-high performance liquid chromatography (Agilent Technologies). Intact mass analysis had an expected mass of 46,797 Da, with the observed MW of 46,794 Da equating to a 0.006% error. Digested gel samples were analyzed using a maxis Impact Ultra-High Resolution QTOF instrument (Bruker Daltonics) coupled to a Dionex 3000 nano-uHPLC (Thermo-Fisher). Data analysis was performed using the SearchGUI/PeptideShaker data analysis software (Compomics). Sequence coverage was obtained for residues 2 to 245 of Aat. Based on sodium dodecyl sulfate-polyacrylamide gel electrophoresis, Aat was estimated to be 90 to 95% of the total protein in the purified sample used for enzyme assays.

Enzyme Assays. The standard protocol for generating the *E. coli* cytoplasmic extract was as follows. A 100 mL culture of *E. coli* BL21 was grown to stationary phase in M9 minimal media (84), collected by centrifugation, suspended in 10 mL phosphate buffer (0.1 M, pH 7.4), and then lysed by sonication on ice. The lysate was centrifuged (12,000 rpm, 20 min) to pellet unbroken cells and debris and then the supernatant was filtered through a 3 K_d MW filter (required multiple filters) to remove proteins but retain metabolites in a batch of extract, which was used for all assays. The standard assay volume was 1.0 mL containing 3 μg purified protein, 4 mM MeA, 4 mM α -ketoglutarate, and 1 mM 5' pyridoxal phosphate and contained in 10 mL sealed serum bottles for tracking CH_4 synthesis as described in *Study Site, Sampling, and Initial Sample Analyses*. Control reactions consisted of the same compositions except the Aat enzyme preparation was inactivated by boiling for 20 min.

Estimating Methane Release from Marine MeA. MeA assimilation rates of 0.005 to 54 $\text{nmol L}^{-1} \text{day}^{-1}$, oxidation rates of 0.09 to 3.43 $\text{nmol L}^{-1} \text{day}^{-1}$, and overall turnover rates of 40 to 57 $\text{nmol L}^{-1} \text{day}^{-1}$ have been reported (62, 67). Assuming MeA conversion to CH_4 is restricted to the top 100 m of the marine water column where the majority of primary productivity occurs, and an average turnover rate of 1 $\text{nmol MeA L}^{-1} \text{day}^{-1}$, the 3.6×10^{19} L of seawater in that volume would result in a MeA \rightarrow CH_4 conversion rate of $1.31 \times 10^{13} \text{ mol yr}^{-1}$, equaling a potential release of 210 Mt $\text{CH}_4 \text{ yr}^{-1}$ ($\text{MW}_{\text{CH}_4} = 16.043 \text{ g} \cdot \text{mol}^{-1}$). Assuming that 0.1, 1, or 10% of marine MeA turnover is due to PLP-Aat homologs capable of the MeA \rightarrow CH_4 , 0.21, 2.1, or 21 Mt $\text{CH}_4 \text{ yr}^{-1}$

would be released. This would account for 1.16, 11.6, and 116% of the global CH₄ emissions from oceanic sources, which are estimated at 18 Mt yr⁻¹ (69).

Data Availability. In addition to the methods described herein, DNA sequence data have been deposited in GenBank (<https://www.ncbi.nlm.nih.gov/>): 1) *aat* gene, [MK170382](https://www.ncbi.nlm.nih.gov/nuclink/MK170382) (85); 2) Illumina libraries, Biosample accession: [SAMN13704654](https://www.ncbi.nlm.nih.gov/biosample/SAMN13704654) (86). All other study data are included in the article and/or *SI Appendix*.

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