Activities of methionine-γ-lyase in the acidophilic archaeon “Ferroplasma acidarmanus” strain fer1

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Abstract: Biogeochemical processes on exposed pyrite ores result in extremely high levels of sulfuric acid at these locations. Acidophiles that thrive in these conditions must overcome significant challenges, including an environment with proton concentrations at pH 3 or below. The role of sulfur metabolism in the archaeon “Ferroplasma acidarmanus” strain fer1’s ability to thrive in this environment was investigated due to its growth-dependent production of methanethiol, a volatile organic sulfur compound. Two putative sequences for methionine-γ-lyase (EC 4.4.1.11), an enzyme known to carry out α,γ-elimination on L-methionine to produce methanethiol, were identified in fer1. Bioinformatic analyses identified a conserved pyridoxal-5′-phosphate (PLP) binding domain and a partially conserved catalytic domain in both putative sequences. Detection of PLP-dependent and L-methionine-dependent production of α-keto compounds and thiol groups in fer1 confirmed the presence of methionine-γ-lyase activity. Further, fer1 lysate was capable of processing related substrates, including D-methionine, L-cysteine, L-cystathionine, and L/D-homocysteine. When the two putative fer1 methionine-γ-lyase gene-coded proteins were expressed in Escherichia coli cells, one sequence demonstrated an ability to carry out α-γ-elimination activity, while the other exhibited γ-replacement activity. These fer1 methionine-γ-lyases also exhibited optimum pH, substrate specificity, and catalytic preferences that are different from methionine-γ-lyases from other organisms. These differences are discussed in the context of molecular phylogeny constructed using a maximum likelihood algorithm based on methionine-γ-lyase sequences from a diverse selection of organisms.

Keywords: acidophiles, archaea, methionine-γ-lyase, L-methionine metabolism, volatile organic sulfur compound

Introduction

Acidophiles are organisms that not only survive, but actually thrive in acidic environments (pH < 3) whereas most other organisms perish. Some of the sites from which acidophiles, including archaea, bacteria, fungi, and protists, have been successfully cultured include hydrothermal sites, bioleaching reactors, and environments rich in pyrite (FeS₂), such as abandoned mines.¹⁻¹⁶ At these mine sites, pyrite is converted to sulfuric acid in the presence of water and oxygen in the following reaction:

\[ \text{FeS}_2 + 14 \text{Fe}^{3+} + 8 \text{H}_2\text{O} \rightarrow 15 \text{Fe}^{2+} + 2 \text{SO}_4^{2-} + 16 \text{H}^+. \]

When chemolithotrophic microbes that utilize Fe²⁺ ions as electron donors are present, this process results in accumulation of high proton concentrations.¹
This biogeochemical process creates extremely acidic conditions, with naturally occurring pH as low as −3.6 being recorded. These highly acidic waters also contain unusually high levels of dissolved metals, with up to 200 g/L reported, creating an acid mine drain that impacts the ecology surrounding these sites.

The highly acidic environment poses two important challenges for acidophiles, ie, how to utilize the readily available proton gradient to generate ATP and how to prevent denaturation of biomacromolecules both on the cell surface and within the cytoplasm. Studies on the archaeon “Ferroplasma acidarmanus” strain fer1 (fer1), which was isolated from an abandoned mine in California, have begun to shed some light on these challenges. This archaeon appears to use tetra-ether-linked lipids as one mechanism of resisting acidification of their cytoplasm. Further, they exhibit resistance to high levels of copper and arsenic, two heavy metals found at acid mine drain sites. Previous work has also tied fer1’s ability to metabolize sulfate from their surroundings to the cellular ATP level, suggesting that sulfate metabolism may play an important role in this organism’s ability to thrive in its habitat. This idea is supported by other data showing that optimal growth of fer1 under heterotrophic conditions requires a minimum of 100 mM sulfate. In contrast, the closest evolutionary relative that requires only 14 mM sulfate for growth, suggesting additional metabolic fates for sulfate in fer1. Radiotracer experiments had previously established that one fate for the sulfate is assimilation into proteins. Another observation concerning the sulfur metabolism of fer1 is the formation of a volatile organic sulfur compound, methanethiol (CH₃SH), during active growth of fer1 cultures. However, the link between assimilatory sulfate reduction and methanethiol production is still missing.

In some organisms, the L-methionine-γ-lyase enzyme (EC 4.4.1.11) catalyzes production of methanethiol from L-methionine. L-Methionine-γ-lyase (MGL) is a multifunctional catalytic enzyme that requires pyridoxal-5’-phosphate (PLP) as a coenzyme. This enzyme typically functions as a tetramer to catalyze the α,γ-elimination or γ-replacement of L-methionine and related substrates, like L-cystathionine. Some forms of MGL can also process L-cysteine using α,β-elimination and β-replacement, as well. So far, MGL activity has been characterized in several organisms, including bacteria, plants, and fungi. In addition to enzymatic characterizations, crystallographic analyses of the MGLs from Pseudomonas putida, Citrobacter freundii, and Entamoeba histolytica have shed light on the structural properties of this enzyme, identifying key amino acid residues in this PLP-dependent enzyme.

Examination of fer1 MGL will provide not only a better understanding of the biological role of sulfate metabolism in this acidophile, but may also offer other novel insights. For example, because extremophiles like fer1 can harbor enzymes possessing properties different from their counterparts in regular microbes, a study of the fer1 MGL may lead to discovery of a new variant of the enzyme. An MGL with new enzymatic properties can offer advantages in several fields. For instance, because MGL is a target for treatment of protist infections and certain types of cancers, having a new source for this enzyme with potentially novel properties may prove useful in therapeutic applications. In addition, a better understanding of the biology of acidophiles regarding methionine and sulfate metabolism could prove beneficial when they are utilized in bioleaching applications. Moreover, volatile organic sulfur compounds are important flavoring compounds in microbiological applications involved with food, such as ripening of different types of cheese, which will make MGL with different pH tolerance a useful tool in designing specific applications. Finally, volatile organic sulfur compounds are also significant contributors to the nucleation process of clouds, which affect the albedo component of global temperature regulation.

In this study, MGL activity in fer1 cells was confirmed, and the activities of putative fer1 MGL orthologs that were heterologously expressed in E. coli were examined. These results paved the way for further analyses of fer1 MGL, which may have a significant impact on our understanding of acidophiles and on applications ranging from cancer therapy to bioleaching.

Methods and materials

Growth and culture

Strain fer1 was cultured as previously described at 35°C in mfer medium. Unless otherwise stated, E. coli and C. freundii strains were propagated using Luria-Bertani medium supplemented with ampicillin 100 µg/mL as needed under standard conditions (35°C either on 1.5% [w/v] agar plates or with 110 rpm shaking in liquid broth culture).

Bioinformatics

Putative MGL coding sequences from fer1 were identified using the P. putida W619 sequence as the query in a BLAST search against the complete fer1 genome. MGL protein sequences from different organisms were retrieved from
GenBank (Table 1). Analyses of fer1 MGL sequences were carried out using ProtParam at the ExPASy site.57 Modeling comparisons were carried out using the Phyre2 algorithm.86 Multiple sequence alignment for protein regions was carried out using the ClustalW2 algorithm.49 A phylogenetic tree for MGL proteins was generated using PhyML, with 300 iterations for bootstrap analysis, and the result was rendered using TreeDyn hosted at the Phylogeny.fr site.50–52

Gene cloning
Forward primers incorporating an NdeI site and reverse primers incorporating an NotI site created for each of the two putative fer1 mgl loci to generate inframe translational fusion for 3′ 6 × His addition in the pET21b vector (EMD Millipore, Billerica, MA, USA) were ordered from Integrated DNA Technology Inc (Coralville, IA, USA). Polymerase chain reaction was carried out following the standard manufacturer’s protocol using the KoD enzyme (EMD Millipore). Purification of DNA fragments from agarose gel was carried out using Wizard® SV Gel and the PCR Clean-up Kit (Promega, Fitchburg, WI, USA). Ligation was carried out using T4 DNA ligase (Promega). The ligated product was transformed into competent BL21(DE3) E. coli cells and transformants were selected for and then propagated. Selected clones were then sequenced for the inserts from both directions in duplicate to confirm sequence integrity.

Cell lysis and protein quantification
To prepare cells for the assays, BL21(DE3) E. coli strains and C. freundii control were grown to mid-log phase and then expressions of cloned proteins were induced by addition of 1 mM isopropyl-β-D-1-thiogalactopyranoside for 3 hours at 20°C with 150 rpm shaking. One hundred mL of overnight cultures were pelleted, rinsed, and then resuspended in 800 µL of 100 mM Tris-HCl (pH 8.0). Lysis was achieved by adding 200 µL 10 × FastBreak® (Promega buffer supplemented with Hal™ Protease Inhibitor Single-Use Cocktail EDTA-Free [Thermo Fisher Scientific, Waltham, MA, USA]) and DNaseI (Promega) and then incubated at 10°C for 30 minutes with 150 rpm shaking. Following lysis, the samples were centrifuged at 13,000×g for 12 minutes at 4°C and the supernatant was aliquoted for assay. For fer1, cell lysis was carried as reported before.23 Total protein was measured using Coomassie Plus Reagent (BioRad, Hercules,

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CA, USA) with a standard curve generated using bovine serum albumin (Sigma-Aldrich, St Louis, MO, USA) protein standards.

Enzyme assays
The activity of MGL was determined using either 3-methyl-2-benzothiazolinone (MBTH Sigma-Aldrich), which reacts with α-keto compounds, or 5,5′-dithio-bis(2-nitrobenzoic acid) (DTNB Sigma-Aldrich), which reacts with thiol groups.53,54 For both assays, cell lysate (40 µL) was combined with 20 µL of 10 mM PLP (Thermo-Fisher) and incubated for 5 minutes at 35°C, followed by addition of reaction substrate and incubation for 25 minutes at 35°C. Substrates, which were dissolved in 50 mM Tris-Cl adjusted to different pH as specified, included: 50 mM L-methionine, 50 mM D-methionine, 50 mM L-cysteine, 5 mM L-cystathionine, and 10 mM DL-homocysteine (final concentrations). All chemicals were purchased from either Sigma-Aldrich or Thermo-Fisher. For the MBTH assays, the reaction was adjusted to a final volume of 1 mL. After incubation with the substrate, 100 µL of 50% (w/v) trichloroacetic acid solution was added and mixed to stop the reactions. Two mL of 1 M sodium acetate (pH 5) and 800 µL of 0.1% (w/v) MBTH were then added and incubated as specified. Production of azine derivatives (extinction coefficient 15.7/mM/cm) indicative of MGL activity were detected from 1 mL of the reaction using spectrophotometric measurement at 320 nm. For DTNB assay, 40 µL of lysate, 5 µL of 1 mM DTT, substrate, and 100 µL of 1 mM DTNB in 2.5 mM sodium acetate solution (pH 2.5) were combined, brought to a final volume of 1 mL, vortexed, and incubated at 25°C for 25 or 30 minutes. Products with an extinction coefficient of 13.6/mM/cm were detected using spectrophotometric measurement at 412 nm. Spectrophotometric measurements were taken using a GenesysII reader (Thermo-Fisher). For both assays, values of samples were corrected by assay blanks containing only the buffer. Enzyme activity (µmol/min) was then calculated using the extinction coefficient and duration of the enzymatic reaction, and corrected for the volume of the reaction actually quantified in the case of the MBTH assay. The specific activity level was calculated by dividing enzyme activity by the amount of protein in the cell lysates (µmol/min/mg).

Statistical analyses
Statistical significance was determined using a two-tailed t-test and reported as the P value. Averages, standard errors, and t-test values were calculated using Microsoft Excel software (Microsoft Corporation, Redmond, WA, USA).

Results
In silico analyses
Similar to the protozoan Entamoeba histolytica, the completed fer1 genome contains two sequences with significant homology to other PLP-requiring enzymes involved in methionine/cysteine metabolism, ie, ZP_05571061 (fer1MGL1) and ZP_05570711 (fer1MGL2, Table 1). Compared with the amino acid sequence of the functionally well characterized MGL from P. putida, fer1MGL1 shares 37% identity and 57% similarity, while fer1MGL2 shares 32% identity and 57% similarity.46 The two fer1 sequences are significantly alike but not identical (38% identity, 54% similarity): fer1MGL1 is predicted to be 42.3 kDa with a pl of 5.34, whereas fer1MGL2 is predicted to be 39.9 kDa with a pl of 6.11.47

The two putative PLP-dependent protein sequences were identified as candidates for methionine-γ-lyase based on ClustalW2 alignment with five MGL sequences that have been previously characterized enzymatically and/or with crystallography, from P. putida (amino acid positions used as reference), C. freundii, Brevibacterium linens, E. histolytica, and Arabidopsis thaliana (Table 1).49 The results show that the four residues known to be critical to binding the PLP cofactor required for enzymatic activity, ie, Met90, Asp187, His207, and Gly215, are present in both fer1MGL1 and fer1MGL2 with the correct spacing between amino acids, suggesting that both proteins from fer1 require PLP as a coenzyme (Figure 1). This prediction is further corroborated by protein-folding modeling using the Phyre2 algorithm, where both sequences exhibited significant structural similarities to PLP-binding regions.48 Of the six residues that are implicated in catalytic activity (Tyr59, Arg61, Tyr114, Cys116, Lys240 and Asp241), both fer1MGL1 and fer1MGL2 showed conservation at the Tyr59 and Arg61 residues. The fer1MGL1 also shares Tyr114, but fer1MGL2 has a Phe114 instead, substituting a polar residue for a nonpolar one, although both contain a ring structure. Noticeably, the Cys116 residue that is critical for catalytic activity in P. putida was not conserved in either of the putative fer1MGL orthologs.55 At that same residue, B. linens and A. thaliana both carry a glycine residue instead of cysteine, whereas fer1MGL1 and fer1MGL2 carry glutamine and arginine, respectively. At the last two of the six catalytic residues, Lys240 and Asp241, fer1MGL1 carries arginine and lysine while fer1MGL2 carries two tandem arginine residues. Compared with P. putida, the two putative fer1 orthologs have conserved a basic residue by substituting Arg240 with...
However, at the second basic residue Arg\textsuperscript{241}, they carry Asp\textsuperscript{241}, which is an acidic residue. Interestingly, the lysine-asparagine doublet is not conserved in \textit{B. linens} or in \textit{A. thaliana}. Given the extent of substitutions and deviations from the conserved catalytic residues, the catalytic characters of these two fer1 enzymes are likely to differ significantly from the ones that have been previously characterized.

To place fer1MGL1 and fer1MGL2 in the evolutionary context of MGL sequences, a phylogenetic analysis using the maximum likelihood method (PhyML) was carried out with an alignment file generated from the ClustalW2 algorithm consisting of organisms from each major evolutionary lineage (Figure 2).\textsuperscript{59,50} While four branch points had low bootstrap values (<50%), most of the rest were well supported.\textsuperscript{56} The results show that although the three fungal sequences examined formed a monophyletic group, the other sequences did not typically align with organismal phylogeny. The crenarchaeal and euryarchaeal sequences were distributed in different branches, intermingling with Gram-positive and Gram-negative bacteria as well as with protozoans. Several archaeal MGL sequences from both crenarchaeal and euryarchaeal members, including fer1MGL1 and fer1MGL2, share higher homology with plant MGLs than they do with bacterial, protozoan, or other archaeal sequences. Notably, fer1MGL1 and fer1MGL2 are not the closest neighbors to each other, reflecting the significant sequence divergence already described. This placement of the two fer1 orthologs is of particular

**Figure 1** ClustalW2 alignment of methionine-\(\gamma\)-lyases protein sequences (see Table 1 for strain abbreviations and accession numbers).

**Notes:** Residues involved in PLP binding are highlighted in gray, whereas residues involved in catalytic reactions are highlighted in black. In both regions, residues that differ from the reference sequence are underscored.

**Figure 2** Phylogenetic relationship among methionine-\(\gamma\)-lyases of different species (see Table 1 for strain abbreviations).

**Notes:** Phylogram was generated using PhyML (maximum likelihood) with 300 bootstrap iterations based on ClustalW2 alignment results. Red arrows indicate branch points with less than 50% support. The colors for different branches correspond to the various classification of the organisms: plants are green, protozoans are orange, fungi are brown, Gram-negative bacteria are pink, Gram-positive bacteria are purple, crenarchaeotic archaea are blue, and euryarchaeotic archaea are black. The scale bar represents branch length in number of amino acid substitutions per residue. Asterisks indicate methionine-\(\gamma\)-lyases that have been confirmed for activity in an enzymatic assay.
interest in comparison with the close placement for the two putative MGLs from the archaeon *Pyrococcus yayanosii* (a euryarchaeote) and from the protozoan *E. histolytica*. In those two organisms, it appears that the two copies of MGL might have resulted from evolutionary divergence following gene duplication, whereas in fer1 the two sequences might have different origins, or it might indicate that they diverged in function much sooner than their counterparts did in the other two organisms.

**Methionine-γ-lyase activities in fer1**

The presence of MGL activity in fer1 was assayed using DTNB with L-methionine as a substrate. Compared with control conditions in which fer1 lysate was denatured at 100°C for 15 minutes, untreated lysate showed significantly higher activity during the assay (*P* < 0.01, Figure 3A). Furthermore, when PLP or the substrate L-methionine was omitted from the assay mixtures, the fer1 cell lysate generated lower levels of thiol products (*P* < 0.05), confirming the PLP-dependence.

![Graph](https://www.dovepress.com/)

**Figure 3** Methionine-γ-lyase (MGL) activity is present in fer1 and varies with pH. (A) Production of free thiol groups from L-methionine by fer1 cell lysate in 5,5′-dithio-bis(2-nitrobenzoic acid) assay was quantified using spectrophotometric measurement at A412. (B) Optimal pH for fer1 MGL activity was assayed by quantifying production of α-keto compounds from L-methionine in 2-methyl-3-benzothiozolinone assays.

*Note:* Results represent data from at least three independent trials and vertical bars represent standard errors.
and substrate-dependence of fer1 lysate, respectively. The effects of pH on the enzymatic activity of MGL were then examined by quantifying the production of α-keto compounds from L-methionine in MBTH assays (Figure 3B). The highest specific activity (0.12 ± 0.02 μmol/min/mg) was detected when the assay buffer was adjusted to pH 4. Furthermore, the substrate specificity of the fer1 lysate was also examined in MBTH assays. Results from three independent trials showed that there was no statistically significant difference in fer1 lysate activity between L-methionine (3.97 ± 0.16 × 10^-2 [μmol/min/mg]), D-methionine [3.87 ± 0.60 × 10^-2 μmol/min/mg], L-cysteine [5.17 ± 0.37 × 10^-2 μmol/min/mg], L-cystathionine [3.53 ± 0.56 × 10^-2 μmol/min/mg], and L(-)-s-homocysteine [6.06 ± 0.97 × 10^-2 μmol/min/mg, P > 0.05]). In contrast, lysate inactivated by boiling showed less activity (0.60 ± 0.14 × 10^-2 μmol/min/mg, P < 0.01) and a positive control using whole cell lysate from C. freundii showed more activity (40.49 ± 4.21 × 10^-2 μmol/min/mg, P < 0.01).

Cloning of fer1 MGLs

Primers for each of the two fer1 MGLs were designed to incorporate NotI (5′ primers) and NdeI (3′ primers) at appropriate positions to create intramolecular translational fusions in the pET21b vector where the coding region is fused with 6× His tag at the carboxyl terminus and is expressed under the control of a promoter that is inducible by isopropyl β-D-1-thiogalactopyranoside. Polymerase chain reactions using a high-fidelity polymerase generated amplicons of appropriate lengths (Figure 4A). To clone the polymerase chain reaction fragments into the pET21b vector, both vector DNA and polymerase chain reaction fragments were first digested by NotI and NdeI simultaneously and then separated by agarose gel electrophoresis. Vector backbone DNA and polymerase chain reaction fragments were then purified from agarose gel slices and mixed for ligation. Chemically competent BL21(DE3) E. coli cells were prepared and used as hosts for ligated products using a heat-shock transformation protocol.27 Plasmids from survivors were isolated and then digested with NotI and NdeI to confirm the length of the inserted polymerase chain reaction product (Figure 4B). The integrity of the cloned inserts was confirmed by sequencing reactions in duplicate from both ends of the cloning sites.

Methionine-γ-lyase activity of fer1MGL1 and fer1MGL2

The MGL activity of heterologously expressed fer1MGL1 and fer1MGL2 was assayed to confirm the functional predictions of these two loci using MBTH assays, which detect the ability of these cell lysates to produce α-keto compounds from L-methionine (Figure 5A). The results showed that the fer1MGL2 cell lysate exhibited activity above control conditions using BL21(DE3) cells without any vectors when assays were carried out at pH values from 2 to 6 (P < 0.05). In comparison, the fer1MGL1 cell lysate did not show significant activity under identical conditions, nor at the other pH values tested (P > 0.05). The validity of the assay was confirmed using C. freundii as a positive control, which showed significant activity at pH values from 5 to 9 (P < 0.05). Further, fer1MGL2 showed a significant increase in specific activity when compared with identical lysates that had been inactivated by boiling at 100°C for 15 minutes (P < 0.05, data not shown).

When the same set of strains were examined for production of thiol groups using DTNB assays adjusted to different pH values, fer1MGL1 exhibited more activity than the BL21(DE3) control at pH 5 (P < 0.05), while fer1MGL2 exhibited the same specific activity as BL21(DE3) at all pH

Figure 4 PCR amplification and cloning of fer1MGL1 and fer1MGL2. (A) PCR amplification of fer1MGL1 (lane 1) and fer1MGL2 (lane 2) produced amplicons of appropriate lengths, 1146 and 1089 bp, respectively, as visualized by 0.8% (w/v) agarose gel electrophoresis. Lane L contains molecular weight marker, where the 4 smallest bands are 500 bp, 1000 bp, 1500 bp, and 2000 bp, with the higher weight bands each at 1000 bp apart. (B) Confirmation of PCR fragment insertion into pET21b was carried out using digestion with NdeI and Ndel (lanes 1 and 3). Notes: Control digestions with identical plasmids using only Ndel were performed as well (lanes 2 and 4). Fragment of correct length was detected for both putative clones of fer1MGL1 (lanes 1 and 2) and fer1MGL2 (lanes 2 and 4). Lane L contains molecular weight marker, where the 4 smallest bands are 500 bp, 1000 bp, 1500 bp, and 2000 bp, with the higher weight bands each at 1000 bp apart.

Abbreviations: MGL, methionine-γ-lyase; PCR, polymerase chain reaction.
values assayed \((P > 0.05, \text{Figure 5B})\). The positive \textit{C. freundii} control showed significant activity at pH values from 5 to 7 \((P < 0.05)\).

**Discussion**

The genome of \textit{F. acidarmanus} strain fer1, which has been completely sequenced, contains only two putative open reading frames with a significant match to PLP-dependent proteins. The automated genome annotation assigned the two loci as cystathionine-\(\gamma\)-lyase (CGL) and a “hypothetical protein” based on sequence homology. However, the degree of amino acid sequence homology between the two sequences (42% identity, 61% similarity for fer1MGL1 and 36% identity, 59% similarity for fer1MGL2) and the previously characterized CGL from the bacterium \textit{Lactobacillus reuteri} (YP_001270900) is comparable with their sequence homology to MGL.58 Further, CGL typically produces hydrogen sulfide from L-cysteine, but actively
Figure 5A). This could be due to the sensitivity of the assay $^\text{2}$-keto acids, the result suggests that fer1MGL1 might have a catalytic preference for α-replacement over α,γ-elimination reactions. This difference in catalytic preference echoes the different substrate preference for the two versions of MGL in E. histolytica and may reflect the divergence in amino acid sequences at the catalytic sites between fer1MGL1 and fer1MGL2 (Figure 1). $^{28}$ In comparison, fer1MGL2 appeared to exhibit activity in MBTH assays but not in DTNB assays (Figure 5A). This could be due to the sensitivity of the assay systems or interference from other cellular components.

Another interesting observation is that, contrary to all the cases where MGL activity had been studied, fer1 cell lysate was able to process D-methionine under DTNB assay conditions. $^{2,26,30,61,62}$ This may reflect the new catalytic residues found in fer1MGL1 and fer1MGL2, which do not match the residues of the other sequences examined in this study. This discovery is in agreement with other cases where fer1 enzymes, such as a DNA repair protein, a DNA helicase, and a DNA ligase, exhibited unusual activity when compared with similar enzymes. $^{37-39}$ Moreover, the pH optimum for the E. coli lysate expressing fer1MGL1 and fer1MGL2 was around pH 5, which is significantly different from the pH optimum reported for the pH 7 value typically reported for other MGLs. $^{29}$ This pH optimum also matches the estimated internal pH for fer1, which is 4.98 ± 0.48. $^{19}$ These new characteristics of fer1 MGLs may be useful in the design of an MGL suitable for therapeutic use in treating certain types of amoebiasis or as anticancer agents against cells that rely on L-methionine for proliferation. $^{40-42}$

The phylogenetic alignment of MGLs from various organisms using a maximum likelihood model showed a surprisingly mixed evolutionary relationship for this enzyme, even when branch points not well supported by bootstrap values (<50%) were disregarded, eg, crenarchaeal and euryarchaeal versions of MGL were found in mixed lineages with each other (Figure 2). This heterogeneity of protein molecular character may be an artifact of using sequence data that have not been functionally confirmed in many of the species examined. Alternatively, this could point to a high frequency of lateral gene transfer events or to the flexibility of many of the PLP-dependent enzymes concerning their substrate-binding capacities.

Regarding substrate specificity, among the MGLs that have been enzymatically characterized, only MGLs from E. histolytica and Trichomonas vaginalis, both protozoans, showed the same pattern of exhibiting comparable levels of activity towards both L-methionine and L-cysteine. In contrast, the other MGLs assayed all showed significantly reduced activity towards L-cysteine when compared with L-methionine. $^{29,30}$ However, when the six catalytic residues and phylogenetic relationships were examined, fer1MGL1 and fer1MGL2 showed less similarity to these protozoan MGLs than they did to plant MGLs. Fortuitously, in a study that examined the role of specific amino acid residues in E. histolytica MGL enzymatic activities, a mutation that changed the Tyr$^{114}$ residue to Phe$^{114}$ was created, mimicking the same substitution found in fer1MGL2. $^{29}$ In the E. histolytica mutation, the MGL activity for the Y114F mutant was significantly reduced in the MBTH assay. In fer1, no activity was detected for fer1MGL1 under MBTH assay conditions, but it did exhibit γ-replacement activity under DTNB assay conditions.

Growth of fer1 requires an unusually high level of sulfate (≥100 μM). $^{21}$ While the sulfate level in fer1 has been shown to correlate with the cellular ATP level, it remains unclear what other roles sulfate may play in the survival of fer1.
Because the $\alpha,\gamma$-elimination of L-methionine carried out by MGL creates methanethiol, ammonia, and 2-oxobutanoate, one positive outcome of MGL activity might be to help stabilize cellular pH by producing ammonia. Alternatively, removal of six protons in the reduction of sulfite to sulfide during sulfate assimilation may serve to remove excess protons from the cytoplasm, with the eventual release of sulfur atoms in methanethiol. Moreover, sulfate assimilation and methanethiol production may serve to maintain a correct balance of redox carriers. With the activity of MGL demonstrated and the connection between sulfate assimilation and methanethiol production established, future studies could further explore the metabolic roles of sulfate and methanethiol in fer1 to elucidate the pH homeostasis mechanisms of this acidophile.

In summary, using whole-cell lysate of fer1, PLP-dependent and L-methionine-dependent production of $\alpha$-keto compounds and thiol groups was demonstrated, thus indicating the presence of MGL in this acidophilic archaeon. Enzymatic activity was also detected in lysates of E. coli cells induced to express cloned fer1 MGL orthologs. These new features of fer1 MGL1 and fer1 MGL2 in their pH optimum, substrate specificity, and catalytic preferences will shed light on the mechanism of sulfur metabolism in this organism, and possibly lead to a better understanding of this organism’s ability to thrive in highly acidic environments. Further experiments using purified fer1 MGL1 and fer1 MGL2 will be required to characterize these enzymes better and to elucidate their possible contribution to other applications, like therapeutic uses and production of volatile organic sulfur compounds. Studies aiming to elucidate the regulatory aspects of these two MGLs will also be valuable in understanding the role of sulfur metabolism in this acidophile. Finally, more work is needed to reconcile the phylogenetic relations of MGLs amongst organisms of various evolutionary lineages.

Disclosure
The authors report no conflicts of interest in this work.

References


