XoxF-Type Methanol Dehydrogenase from the Anaerobic Methanotroph “Candidatus Methylomirabilis oxyfera”

Ming L. Wu,*† Hans J. C. T. Wessels,‡ Arjan Pol,§ Huub J. M. Op den Camp,¶ Mike S. M. Jetten,¶ Laura van Niftrik,* Jan T. Keltjens*‡

Department of Microbiology, Radboud University Nijmegen, Institute for Water and Wetland Research, Nijmegen, The Netherlands;† Nijmegen Centre for Mitochondrial Disorders, Centre for Proteomics, Glycomics and Metabolomics, Laboratory of Genetic, Endocrine, and Metabolic Disease, Department of Laboratory Medicine, Radboud University Medical Centre, Nijmegen, The Netherlands

The bacterium “Candidatus Methylomirabilis oxyfera” is a recently discovered player in the methane cycle. This organism, which is the first representative of the novel division of NC10 bacteria with a known physiological function, derives its energy from growth from the anaerobic oxidation of methane coupled to the reduction of nitrate or nitrite into dinitrogen gas (1, 2). “Ca. Methylomirabilis oxyfera” is an extraordinary organism since methane oxidation to CO2 proceeds via methanol, formaldehyde, and formate. In this pathway, the first step, activation of methane into methanol, requires oxygen and this oxygen is produced by “Ca. Methylomirabilis oxyfera” itself in concert with nitrite reduction (2). The second step is the reduction of methanol. This reaction is catalyzed by methanol dehydrogenase (MDH). MDH is a key enzyme widely distributed among aerobic methane-oxidizing (“methylotrophic”) and methyl group-oxidizing (“methylophilic”) microorganisms. MDHs fall into different classes. Gram-positive microorganisms harbor nicotinoprotein MDHs in their cytoplasm that use NAD(P)H as the electron acceptor for methanol oxidation (3, 4). In Gram-negative bacteria, a phylogenetically unrelated protein is present in the periplasm that possesses pyrroloquinoline quinone (PQQ) as the catalytic center (5–7). PQQ-containing MDHs are members of a quinoprotein family that are able to oxidize a broad range of alcohol and aldehyde substrates, and each member is more or less tuned for a specific substrate (5–8).

Within genomes, a large variety of protein sequences have been annotated as (putative) quinoprotein MDHs. Functionally and structurally best understood are the so-called MxaFI MDHs. These enzymes are well characterized by the resolution of their atomic structures of six different bacterial species (9–16). MxaFI MDHs are heterotetrameric (α2β2) enzymes composed of two large (α, MxaF) and two small (β, MxaI) subunits. Each large subunit contains one noncovalently bound PQQ prosthetic group and one Ca2⁺ ion at its active site, both of which are essential for enzyme activity (6). The highly basic small subunit, which is not found in other quinoproteins, tightly wraps against the large subunit, but its function is not fully clear. Active MxaF MDHs, thus lacking the β subunit, have been purified before, suggesting that this subunit is not always essential for enzyme activity per se (17, 18). The functional expression of MxaFI MDH requires the action of up to 30 genes. In Methylobacterium and Methylococcus species, these genes are spread over five gene clusters (mxa, mxd, mxc, pqqABCDE, and pqqFG) (19–21). In the mxa cluster, mxaF and mxaI are the structural genes. The mxaG gene codes for cytochrome c1, the cognate electron acceptor during methanol oxidation. Occasionally, the mxaI gene product has been observed as an additional subunit (22, 23), but most of the MxaFI MDH prep-
rations described to date lack this protein, even though the mxaF and mxaI genes are commonly linked to each other in genomes (8). The mxaACKLD gene products are essential for Ca\(^{2+}\) insertion into the apoprotein (19, 24), whereas the products of the mxaRSAHI genes are thought to play a role in further MDH maturation (25–27). MxaB is a DNA-binding protein involved in transcriptional regulation, as are the two component regulators encoded by the mxb and mxc clusters (28). The gene products of the pqqABCDE and pqqEF clusters are involved in PQQ biosynthesis for which the 23- to 24-amino-acid (aa) gene product of pqqA is the proposed precursor of PQQ (29–31).

Genome sequencing projects revealed the presence of MxaF homologs, termed XoxF proteins, that showed, at most, 50% sequence identity to known MxaFs (32–34). XoxF genes can be detected in many genomes, not only of methanotrophic and methylo trophic species, but also of microorganisms that have not been implicated in such a lifestyle (8, 34). Phylogenetic analysis divides XoxF proteins into at least five different clades (XoxF1 to XoxF5) (8, 34). Despite their widespread occurrence, the function of XoxF proteins has been enigmatic for some time. Unlike mxaF, xoxF genes were hardly expressed during growth under laboratory conditions and their deletion did not result in a clear phenotype (32). In striking contrast, xoxF genes were highly expressed in the plant phyllosphere (35) and by communities of nutrient-limited coastal ocean waters (36). Certain methanotrophs, such as the Verrucomicrobia (37), lack genes coding for an MxaF MDH, and an XoxF protein would be the only candidate for methanol oxidation. In agreement with this, the deletion of xoxF from Rhodobacter sphaeroides resulted in the loss of this phototroph’s ability to use methanol for photorespiration and aerobic respiration (38). Moreover, the purification of XoxF proteins from different bacterial species established their function as MDHs (39–43). These studies also shed light on their elusive nature. The proteins were specifically induced in the presence of rare earth elements (REEs) like La\(^{3+}\) or Ce\(^{4+}\) (40–42). The recent resolution of the crystal structure of the XoxF MDH from the Verrucomicrobia phylum member Methylocystis fumaricolum SolV showed that this REE took the position near the PQQ catalytic site usually occupied by calcium in MxaF MDHs (43). Importantly, all of the XoxF MDHs described thus far are homodimeric proteins lacking a small subunit.

The genome of “Ca. Methylophilobacter oxyfera” codes for three MxaF homologs that have been annotated as MxaF1 to MxaF3 and that are encoded by one long gene cluster (2). However, key genes that are encoded by this cluster lack genes coding for an MxaF MDH, and two different XoxF MDHs. To address the question of which of these proteins is functionally expressed, we purified MDH from the organism. Curiously, the only active MDH that could be isolated was a heterotetrameric enzyme composed of two XoxF1-type large subunits and two small subunits, and it contained PQQ as its prosthetic group. Using specific antibodies raised against the XoxF1 large subunit, we localized the enzyme to the periplasm of “Ca. Methylophilobacter oxyfera” cells.

**MATERIALS AND METHODS**

“Ca. Methylophilobacter oxyfera” enrichment culture. “Ca. Methylophilobacter oxyfera” strain Ooij was enriched from a sediment sample taken from a ditch draining agricultural land in the Ooij polder, a floodplain of the River Rhine in The Netherlands, by the sequencing batch reactor technology (45). Enrichment, growth, and culture maintenance in a 15-liter bioreactor were performed as described before (1, 2, 46–49). Growth took place in an atmosphere of 95% CH\(_4\)-5% CO\(_2\) (vol/vol) with nitrile as the electron acceptor. The trace elements present in the mineral medium were as specified in reference 48 and lacked any additional REEs. Here, we also added boiled cell extract from the methanotroph *M. fumaricolum* SolV (43, 50). The addition of the extract (10 ml weekly) resulted in an immediate increase in the methane and nitrite conversion rates; cellular activities had increased no less than 10-fold after 2 months. In the biomass that was collected from the enrichment culture, which was also used in parallel studies (2, 46–49), “Ca. Methylophilobacter oxyfera” made up about 80% of the population, as shown in these studies by fluorescent in situ hybridization and metagenome analysis. The residual community (about 20%) was highly diverse and evenly distributed.

**Sequence and expression analyses.** Sequences of the “Ca. Methylophilobacter oxyfera” MDH gene clusters and homologues of PQQ genes were retrieved from the NCBI genome database (BioProject accession number PRJNA161901). Sequences of MDH and PQQ gene clusters of *M. extorquens* AM1 were retrieved from GenBank and NCBI databases. Protein sequence analyses and comparisons used the BLAST (http://blast.ncbi.nlm.nih.gov/Blast.cgi), ClustalW2 (http://www.ebi.ac.uk/Tools/msa/clustalw2/), and HHpred homology detection and structure prediction tools (http://toolkit.tuebingen.mpg.de/hhpred) tools. The SignalP 4.1 program (http://www.cbs.dtu.dk/services/SignalP/) (51) was used to predict N-terminal leader sequences and cleavage sites. Levels of transcription are expressed as transcriptome coverage by short-read Illumina sequences (32 nucleotides [nt]) of reverse-transcribed RNA obtained from the Sequence Read Archive (BioProject accession number SRR0227482) (2).

**Enzyme purification.** Approximately 26 g (wet weight) of cells from the “Ca. Methylophilobacter oxyfera” enrichment culture were harvested by centrifugation (6,000 \(\times\) g, 10 min, 4°C). This and all subsequent steps were performed aerobically. Purification was monitored by measuring MDH activity immediately after each step. Pelleted cells were washed twice in 20 mM potassium phosphate (KP\(_2\)) buffer (pH 7.0) and resuspended in 20 ml of KP buffer containing 50 mM sodium pyrophosphate and a few grains of solid DNase. Cells were broken by sonication. Cell debris was removed by centrifugation (6,000 \(\times\) g, 10 min, 4°C), and the supernatant (1.2 mg of protein ml\(^{-1}\)) was collected as cell extract. The fraction containing soluble proteins was obtained as the supernatant after ultracentrifugation of the cell extract (143,000 \(\times\) g, 60 min, 4°C). This clear supernatant, saturated ammonium sulfate was added dropwise at 4°C. MDH activity precipitated between 2.0 and 2.8 M (NH\(_4\))\(_2\)SO\(_4\). Following centrifugation (6,000 \(\times\) g, 15 min, 4°C), the precipitated protein fraction was resuspended in 10 ml of 20 mM KP buffer and applied to a column (1 ml) packed with CHT Ceramic Hydroxyapatite (Bio-Rad, Hercules, CA). The column was equilibrated with washing with 5 column volumes of KP buffer. Elution was performed with a linear phosphate (pH 7.0) gradient (20 to 500 mM at a flow rate of 1 ml min\(^{-1}\)). The only fraction containing MDH activity was eluted at 125 mM phosphate. This fraction was concentrated with a spin column with a 5-kDa cutoff (Sartorius Stedim Biotech, Aubagne, France). Fast protein liquid chromatography (FPLC) column purification was performed at room temperature with an AKTA purifier (GE Healthcare, Uppsala, Sweden).

Protein concentrations were determined by the method of Bradford (52) with the Bio-Rad protein assay kit (Bio-Rad, Hercules, CA) with bovine serum albumin (BSA) as the standard. Protein fractions obtained after each step and the purified MDH were analyzed by SDS and native PAGE. Benchmark protein standards [LC0725 NativeMark [Invitrogen Life Technologies, Bleswijk, The Netherlands] for native PAGE, SM0761 prestained PAGE ruler [Fermentas-Thermo Fisher Scientific, Waltham, MA] for SDS-PAGE] were used to estimate molecular masses. Gels were stained with Coomassie brilliant blue G250 (Bio-Rad).

**Enzyme assays.** MDH activity was measured spectrophotometrically at 30°C by monitoring the methanol-dependent and phenazine metho-
sulfate (PMS)-mediated reduction of 2,6-dichlorophenol indophenol (DCPIP) at 600 nm ($\varepsilon_{600}$ of 21,500 M$^{-1}$ cm$^{-1}$) essentially as described by Antony and Zatman (53) with 100 mM Tris-HCl (pH 8.0) as the buffer and increased concentrations of methanol (20 mM) and NH$_4$Cl (45 mM). Reaction mixtures (0.5 ml) contained 1 mM KCN to suppress the reoxidation of reduced DCPIP. Enzyme kinetic parameters were assessed by nonlinear regression analysis with the Origin 8.5.1 program (OriginLab Corporation, Northampton, MA) by using Michaelis-Menten kinetics.

Analytical ultracentrifugation. Analytical equilibrium ultracentrifugation was performed at 20°C in a Beckman XL-I ProteomeLab ultracentrifuge (Beckman Coulter, Fullerton, CA) equipped with an An-60 Ti rotor and cells with a 1.2-cm path length. Prior to ultracentrifugation, the protein was equilibrated in 25 mM HEPES-KOH buffer (pH 7.5) containing 25 mM KCl and 1 mM methanol and adjusted to an A$_{280}$ (1 cm) of 0.75. The speed used was 30,000 rpm. Equilibrium data were evaluated for 300 scans with the SEDFIT program (54) and assuming a fractional ratio ($f_{i}/f_{o}$) of 1.05 for a slightly ellipsoid protein.

UV-visible light absorption spectroscopy and ICP-MS. The absorption spectrum of the purified enzyme was recorded in a cuvette with a 1-cm path length at room temperature on a Cary 50 spectrophotometer. The ICR cell was programmed to acquire selected ion monitoring (SIR) scans of 314 to 344 m/z against the [M + H]$^+$, [M + H$^+$] + 2, and [M + 2H]$^+$ peaks for determination of molecular masses. To verify the presence of PQQ, tandem MS was employed. A protein mixture was loaded at a flow rate of 1.2 l/min directly onto the analytical mass spectrometer (LTQ FT Ultra; Thermo Fisher Scientific). Samples were subjected to tryptic in-gel digestion according to Wilm et al. (56). Peptides were extracted and prepared for matrix-assisted laser desorption ionization–time of flight (MALDI-TOF) MS as described by Farhoud et al. (57). MS of the tryptic digests was performed on a Bruker Biflex III spectrometer (Bruker Daltonics, Fremont, CA) operated in reflectron mode. Spectra were analyzed with the Mascot Peptide Mass Fingerprint search program (Matrix Science, London, United Kingdom) against the “Ca. Methylovorans oxyfera” database, with oxidation (M) as variable modification, 0.2 Da peptide tolerance, and a maximum of one missed cleavage. Molecular masses of the native large and small subunits were determined by MALDI-TOF MS operating in the linear mode. These analyses of the is-asolated enzyme were performed by using the [M + H]$^+$, [2M + H]$^+$, [M + 2H]$^+$, and [M + 3H]$^+$ peaks for determination of molecular masses.

RESULTS
Genomic organization of three MDH systems of “Ca. Methylovorans oxyfera.” The genome of “Ca. Methylovorans oxyfera” codes for three different MDH systems (MDH-1, MDH-2, and MDH-3) that are located on the same strand of a cluster of partly overlapping genes (DAMO_0112 to DAMO_0138) (Fig. 1). The cluster is made up of three subclusters. Each subcluster is preceded by genes encoding proteins (DAMO_0112, DAMO_0124, and DAMO_0134) showing 40 to 74% amino acid sequence identity both to each other and to the large subunits (MxaF and XoxF) of well-defined MxaF and XoxF MDHs (see next). In all three subclusters, genes coding for the putative MDH large subunits are linked to mxaF and mxaG homologs (termed xoxF and xoxG in the
case of XoxF systems). Herein, distinct MxaG/XoxG proteins with the typical CXCH motif for heme c binding would represent the cognate physiological electron acceptor for methanol oxidation, cytochrome \( c_1 \). The function of the MxaI/XoxI-like proteins (DAMO_0113, DAMO_0125, DAMO_0136) is not known, but sequence analysis identified it as a member of the family 3 extra-cellular solute-binding proteins (COG0834, pfam13533), suggesting a role in the binding of methanol or the release of the toxic reaction product formaldehyde. Only the MDH-1 subcluster harbors a gene (DAMO_0115) coding for the small (\( \beta \)) subunit of canonical heterotetrameric MxaFI MDHs. Similarly, only the MDH-1 subcluster comprises the nearly complete set of genes coding for proteins involved in PQP and \( \text{Ca}^{2+} \) insertion (\( \text{mxa}^{	ext{ACKLD}} \); DAMO_0118 to DAMO_0121) and in MDH maturation (\( \text{mxaRSE} \); DAMO_0116 to DAMO_0117, DAMO_0122) (Fig. 1). In “Ca. Methylomirabilis oxyfera,” the \text{mxaL} and \text{mxaD} genes are fused (DAMO_0121). The gene order in the MDH-1 subcluster is the same as that found in \text{M. extorquens} \( \text{a}^\) (19) and in Mxa systems in other organisms (8). However, homologs of \text{mxaH} and the regulatory \text{mxa}AB are absent, which is not unusual (8).

MDH-2 and MDH-3 subclusters lack nearly all of the genes coding for \( \text{Ca}^{2+} \) insertion and maturation proteins, which seems to be a common property of XoxF systems (8). In this respect, the presence of an \text{MxaE} homolog (DAMO_0128) in the MDH-2 sub-cluster is remarkable. The two subclusters are separated by five genes (DAMO_0129 to DAMO_0133) (Fig. 1), of which DAMO_0129 and DAMO_0131 encode putative TonB-dependent and TonB-like transporter proteins, respectively. It is interesting that homologs of DAMO_0129 and DAMO_0131 are widely detected in genomes of XoxF-containing methylotrophs (data not shown). DAMO_0132 and DAMO_0133 code for “Ca. Methylomirabilis oxyfera”-specific exported surface proteins that are structurally characterized by \( \beta \)-propeller strands. Deep RNA sequencing indicated that the DAMO_0130 coding region is one of the most highly expressed parts of the genome (Fig. 1), but it remains to be established whether its transcript is translated into an unknown protein (as annotated) or it is a small noncoding RNA that has some regulatory function.

As described below, all of the amino acids involved in the binding of the PQP prosthetic group are conserved in all three MDH large subunits from “Ca. Methylomirabilis oxyfera.” Moreover, in this work, the presence of this cofactor in one of these MDHs was confirmed by direct purification. Therefore, it was puzzling that the genome of the organism possessed only three of the seven or eight genes required for \( \text{Ca}^{2+} \) biosynthesis (44), i.e., DAMO_0005 (\text{ppqF}), DAMO_0004 (\text{ppqG}), and DAMO_0982 (\text{ppqE}) (2, 44). However, reexamination of the gene products revealed only low sequence similarity to the particular PQP biosynthesis enzymes. Like validated PPQ (760 aa) (30), DAMO_0005 (427 aa) and DAMO_0004 (448 aa) belong to the \text{Zn}-dependent M16 peptidase superfamily. DAMO_0982 (371 aa) and PPQE (~380 aa) are members of the radical S-adenosylmethionine superfamily. Significant differences in protein length and low sequence similarities (<20%) with respect to known PQP biosynthesis genes suggest other functions for these three “Ca. Methylomirabilis oxyfera” proteins, if they are expressed. Hence, “Ca. Methylomirabilis oxyfera” seems to be devoid of PQP biosynthesis machinery, at least as far as we know.

**Sequence analysis of the MDH large and small subunits of “Ca. Methylomirabilis oxyfera.”** The sequences of the three MDH large subunits (DAMO_0112, DAMO_0124, and DAMO_0134) are 42 to 50% identical, and nearly all of the amino acids that had been implicated in MDH crystal structures with the binding of PQP and metal (calcium) prosthetic groups are conserved in these subunits (see Fig. S1A in the supplemental material). Major differences included deletions and/or insertions of small amino acid stretches. More-detailed sequence comparison and phylogenetic analysis assigned these large subunits to three different phylogenetic clades (see Fig. S2 in the supplemental material). DAMO_0112 was affiliated with the MxaF MDHs. At the amino acid level, DAMO_0112 (referred to here as MxaF\( _{\text{Mo}} \)) showed 69 to 74% sequence identity to MxaF MDHs with validated function and known crystal structures. According to the numbering system of Chistoserdova (34), DAMO_0124 and DAMO_0134 are members of the XoxF1 and XoxF2 MDHs, respectively. XoxF1 from “Ca. Methylomirabilis oxyfera” (DAMO_0124; XoxF1\( _{\text{Mo}} \)) showed the highest sequence identity (67 to 74%) to putative MDHs from methanotrophic \text{Methylotenera}, \text{Methyloferula}, and \text{Methyllocella} species, as well as \text{Hyphomicrobium} and \text{Xanthomonas} representatives. XoxF2 from “Ca. Methylomirabilis oxyfera” (DAMO_0134; XoxF2\( _{\text{Mo}} \)) showed the highest sequence identity (62 to 63%) to tentative MDHs from the thermoacidophilic methanotrophic \text{Verrucomicrobia} (37), including the XoxF-type MDH from \text{M. fumarolicum} SoV (XoxF\( _{\text{Mo}} \)), the crystal structure of which has been resolved (43). A particular feature of XoxF\( _{\text{Mo}} \) is the presence of a lanthanide REE (La\(^{3+}\), Ce\(^{3+}\)) at its catalytic site instead of \( \text{Ca}^{2+} \). This REE is coordinated in close proximity to PQP by three highly conserved amino acids (Glu172, Arg256, and Asp299; numbering according to that of processed XoxF\( _{\text{Mo}} \)) that are also involved in the coordination of \( \text{Ca}^{2+} \) (8, 43). However, the proper coordination of the REE requires one more amino acid, Asp301. Besides other sequence characteristics, the presence of Asp301 is a highly diagnostic property of XoxF MDHs (8). Next, two amino acid substitutions (Pro259→Thr, Ala171→Gly) are observed to accommodate the larger REE in the XoxFM\( _{\text{Mo}} \) structure (43). Sequence comparison showed that Asp301 was present in both XoxF1\( _{\text{Mo}} \) and XoxF2\( _{\text{Mo}} \) (see Fig. S1A in the supplemental material). In addition, XoxF2\( _{\text{Mo}} \) displayed the same proline-to-threonine and alanine-to-glycine substitutions seen in XoxF\( _{\text{Mo}} \) but these substitutions were different in XoxF1\( _{\text{Mo}} \) (Pro→Asp, Thr→His) (see Fig. S1A). These observations might indicate that both XoxF MDHs from “Ca. Methylomirabilis oxyfera” possess a REE instead of calcium.

As mentioned before, only the MDH-1 subcluster contained a gene coding for a small subunit, namely, DAMO_0115 (Fig. 1). DAMO_0115 was translated as a polypeptide (MxaI\( _{\text{Mo}} \)) of 94 aa. SignalP analysis (51) suggested a 25-aa leader sequence, pointing to the translocation of the processed protein across the cytoplasmic membrane. The processed protein (69 aa; theoretical molecular mass, 8,075.11 Da) showed 61 to 65% sequence identity to the small subunit of MDHs with known crystal structures (see Fig. S1B in the supplemental material). In these structures, the small subunit tightly binds to the large subunit through a conserved set of large-subunit amino acids (see Fig. S1A). To quite an extent, these amino acids are also conserved in XoxFs, even though the latter may not contain such a small subunit. Like MxaI\( _{\text{Mo}} \), all three MDH large subunits from “Ca. Methylomirabilis oxyfera” were predicted to contain N-terminal leader sequences, again suggesting a periplasmic localization of the processed enzymes. After cleavage of the N-terminal 29 or 30 aa, this would result MxaF\( _{\text{Mo}} \)
XoxF1Mo, and XoxF2Mo, apoprotein molecular masses of 63,526, 67,660, and 63,713 Da, respectively.

Deep RNA sequencing (2) indicated that mxaF was most highly expressed at the mRNA level (transcriptome coverage: mxaF1Mo, 62-fold; xoxF1Mo, 14-fold; xoxF2Mo, 6-fold) (Fig. 1). Strikingly, XoxF1Mo was most highly expressed in the proteome of “Ca. Methylomirabilis oxyfera” strain Ooij, which was used in the present study, whereas XoxF2Mo was most abundant in the proteome of closely related “Ca. Methylomirabilis oxyfera” strain Twente (2), despite the fact that both strains were enriched and cultured under similar conditions. The above considerations raised the questions of which of the three MDHs is (are) functionally expressed, whether their large subunit contains PQQ or an alternative cofactor, which metal is present at the catalytic site, and where the MDH is localized. To address these questions, we purified the MDH from “Ca. Methylomirabilis oxyfera” and assessed its cellular localization by immunogold labeling.

Purification of a heterotetrameric XoxF MDH from “Ca. Methylomirabilis oxyfera.” MDH was purified with a 32% yield by the two-step purification procedure summarized in Table 1. The preparation obtained after the hydroxyapatite step represented the predominant MDH activity (>95%); a few other fractions displayed negligible activity and were not further investigated. After ultracentrifugation of the cell extract, 93% of the total activity was recovered in the supernatant. Hence, MDH is soluble or only loosely membrane associated. The purification factor (8.4-fold) suggested that MDH is a major protein. The purified enzyme catalyzed methanol oxidation by simple Michaelis-Menten kinetics with apparent Vmax and Km values of 10 μmol min⁻¹ mg⁻¹ of protein⁻¹ and 17 μM, respectively. The presence of ammonium strongly stimulated its activity; in the absence of ammonium, it was 10-fold less active.

On native PAGE, purified MDH showed one single band with an apparent molecular mass of approximately 260 kDa (Fig. 2A).

<table>
<thead>
<tr>
<th>Purification step</th>
<th>Total protein (mg)</th>
<th>Sp act (μmol min⁻¹ mg⁻¹)</th>
<th>Total activity (μmol min⁻¹)</th>
<th>Yield (%), estimated by SDS-PAGE</th>
</tr>
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<tbody>
<tr>
<td>Cell extract</td>
<td>24.5</td>
<td>1.10 ± 0.06</td>
<td>26.4</td>
<td>100</td>
</tr>
<tr>
<td>Soluble fraction</td>
<td>19.7</td>
<td>0.97 ± 0.05</td>
<td>19.2</td>
<td>73</td>
</tr>
<tr>
<td>Membranes</td>
<td>3.4</td>
<td>0.53 ± 0.03</td>
<td>1.8</td>
<td>7</td>
</tr>
<tr>
<td>(NH₄)₂SO₄</td>
<td>5.7</td>
<td>2.94 ± 0.23</td>
<td>16.9</td>
<td>64</td>
</tr>
<tr>
<td>Hydroxyapatite</td>
<td>0.9</td>
<td>9.66 ± 1.52</td>
<td>8.4</td>
<td>32</td>
</tr>
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</table>

MALDI-TOF MS gave a molecular mass of 67.3 ± 0.27 kDa (n = 7) for the large subunit (see Fig. S3), which agrees with both that estimated by SDS-PAGE and that expected for N-terminally cleaved XoxF1Mo (DAMO_0124; 67.66 kDa). It should be noted that the molecular masses of the other two large subunits are ~4 kDa less (MxaF_Mo, 63,526 Da; XoxF2Mo, 63,713 Da), differences that are incompatible with MS. Upon analytical equilibrium ultracentrifugation of native MDH, the dominant 280-nm-absorbing band could be modeled (SEDFIT) to an ~150-kDa protein (Fig. 2D), which is clearly less than that observed by native PAGE. This molecular mass of 150 kDa is consistent with an αβ₂ protein composed of two 67-kDa large subunits and two 8-kDa small subunits. Besides this major band, three more bands were obtained during analytical ultracentrifugation of the apparently pure protein preparation (Fig. 2B). These bands sedimented at ~67, ~240, and ~310 kDa. The molecular mass of the first band (67 kDa) again agreed with that expected for XoxF1Mo. Both high-molecular-mass bands might represent higher aggregates (αβ₂, αβ₂). Together, the observations concluded that the MDH purified from “Ca. Methylomirabilis oxyfera” was predominantly a heterotetrameric (αβ₂) enzyme composed of two MxaF_Mo small subunits and two XoxF1Mo large subunits.

The XoxF MDH from “Ca. Methylomirabilis oxyfera” contains PQQ. The UV-visible light absorption spectrum of the purified MDH from “Ca. Methylomirabilis oxyfera” exhibited the characteristics of a quinoprotein (Fig. 3), having an absorption maximum at 345 to 350 nm and a wide shoulder at 375 to 400 nm (60, 61). The spectrum of the enzyme (6.2 mg ml⁻¹) had an absorbance at 342 nm of 0.91. Assuming a PQQ molar coefficient of absorption at 342 nm of 9,620 M⁻¹ cm⁻¹ (55), the quinone concentration was 95 μM, which indicated that the protein bound PQQ in an about 1:1 stoichiometry. The prominent absorption peak around 345 nm was also indicative of the presence of a metal ion in the active site of the enzyme (62, 63). Incubation of the enzyme preparation with EGTA had no effect on the overall spectrum of the enzyme, indicating that the metal would be tightly bound. To assess the nature of the metal, ICP-MS was used as described before (43). The analysis revealed the presence of calcium only; rare earth metals were below the limit of detection (<0.1 mol%). Since column fractions lacking MDH contained calcium as well, presumably derived from hydroxyapatite [Ca₁₀(PO₄)₆OH] used for purification, it was not possible to determine the content of this compound in the protein.

The presence of PQQ in the purified MDH from “Ca. Methy- lomirabilis oxyfera” was confirmed by MS with both precursor ion and fragmentation level data. The acquired precursor ion spectra from the PQQ standard and the purified MDH fully compared the simulated spectrum of PQQ. The monoisotopic mass of the [M+H]-PQQ ion from both the standard MDH (m/z 329.0053; Δ = 4.0 ppm) and the purified MDH (m/z 329.0052; Δ = 3.6 ppm) accurately matched the calculated (deprotonated) monoisotopic mass of PQQ (m/z 329.0040) (see Fig. S4A in the supplemental material). Moreover, the relative abundance of the 13C isotope peaks in precursor spectra from both the PQQ standard (17%) and the purified MDH (16%) was in good agreement with the simulated spectrum (15%). The isotope abundance spectrum was simulated according to de Hoffmann and Stroobant (64). The presence of PQQ in the purified MDH was also confirmed by comparing the collision-induced dissociation fragmenta-

TABLE 1 Purification of MDH from “Ca. Methylomirabilis oxyfera” enrichment culture

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<td>(NH₄)₂SO₄</td>
<td>5.7</td>
<td>2.94 ± 0.23</td>
<td>16.9</td>
<td>64</td>
</tr>
<tr>
<td>Hydroxyapatite</td>
<td>0.9</td>
<td>9.66 ± 1.52</td>
<td>8.4</td>
<td>32</td>
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tion spectra of m/z 329 from the PQQ standard and the purified enzyme (see Fig. S4B). The two spectra were nearly identical and showed the presence of two abundant fragment ions corresponding to the loss of CO₂ (m/z 285 [M-H-CO₂]⁻) and 2CO₂ (m/z 241 [M-H-2CO₂]⁻) from the precursor ion. The findings agreed with those of Noji et al. (65), except that in neither the standard nor the purified enzyme was a third diagnostic fragment ion (m/z 197 [M-H-3CO₂]⁻) detectable, which was probably due to different instrument characteristics and/or conditions.

Cellular localization of the XoxF MDH in “Ca. Methylomirabilis oxyfera.” The N-terminal signal sequences in XoxF1_Mo suggested a periplasmic localization of the processed protein (see Fig. S1A in the supplemental material). To investigate this, we generated primary antiserum with a synthetic peptide targeting a sequence of XoxF1 (α-XoxF1) that was specific for this protein (see Fig. S1A). The specificity of the derived antiserum was con-

FIG 2 PAGE, immunoblotting, and analytical ultracentrifugation of MDH from “Ca. Methylomirabilis oxyfera.” (A) Native 10% PAGE of purified MDH (6 μg). (B) SDS–15% PAGE of the purified MDH (4 μg). The upper band corresponds to XoxF1, and lower band corresponds to MxaI. Marker proteins and their corresponding molecular masses are shown in the left lanes of panels A to C. (C) Immunoblot analysis of the affinity-purified antiserum (α-XoxF1) directed against XoxF1 of “Ca. Methylomirabilis oxyfera.” For SDS–10% PAGE, gels were loaded with cell extract from “Ca. Methylomirabilis oxyfera” (30 μg of protein) or with purified MDH (10 μg) and blotted onto a nitrocellulose membrane. Lanes: 1, marker proteins; 2, cell extract blot incubated with only secondary antiserum; 3, cell extract blot incubated with α-XoxF1 antiserum; 4, purified MDH blot incubated with α-XoxF1 antiserum. The expected target size (~67 kDa) is indicated by the arrow. The values to the left are molecular sizes in kilodaltons. (D) Analytical ultracentrifugation of purified MDH. Analytical ultracentrifugation was performed as described in Materials and Methods, and equilibrium data were fitted by the SEDFIT program (54). Concentrations are expressed in arbitrary units (AU) of absorbance at 280 nm.

FIG 3 UV-visible light absorption spectrum of purified MDH of “Ca. Methylomirabilis oxyfera.” The spectrum was recorded in 150 mM phosphate buffer (pH 7.0) at 30°C. The protein concentration was 6.2 mg ml⁻¹.
The genome of the anaerobic methanotroph “Ca. Methylomirabilis oxyfera” codes for three different MDH systems that are localized in one large gene cluster, which is highly unusual: one MxaFI MDH and two XoxF MDHs (Fig. 1). Here, we purified the predominant MDH that is functionally expressed under the growth conditions used (Fig. 2 and Table 1). The enzyme was composed of two small and two large subunits, as is common among MxaFI MDHs (5–8). MALDI-TOF MS identified the small subunit as the N-terminally cleaved gene product of DAMO_0115 (Mxalmo). Five lines of evidence (SDS-PAGE, MALDI-TOF MS of the native and trypsin-cleaved protein, analytical ultracentrifugation, antibody specificity) consistently indicated that the large subunit with a size of 67 kDa was XoxF1Mo (DAMO_0124). Immunogold labeling localized the protein in the periplasm (Fig. 4), as is also known for MDHs from other methanotrophs and methyloptrophs. In the established (PMS and DCPIP) dye-coupled assay (6, 53), “Ca. Methylomirabilis oxyfera” MDH catalyzed the oxidation of methanol with high affinity (Km of 17 μM) and a high Vmax of 10 μmol min⁻¹ mg⁻¹. The apparent Km for methanol is comparable to those of well-investigated MxaFI MDHs (2 to 20 μM), while the Vmax of the “Ca. Methylomirabilis oxyfera” enzyme was substantially higher than those of known MxaFI MDHs (0.8 to 1.0 μmol min⁻¹ mg⁻¹) (60, 63, 66). In addition, MDH was a major enzyme in “Ca. Methylomirabilis oxyfera.” Taking into account that the very slowly growing bacterium “Ca. Methylomirabilis oxyfera” oxidizes methane at an extremely low rate (1.7 nmol min⁻¹ mg⁻¹ of protein⁻¹) (2, 46), the abundance of MDH and its favorable catalytic efficiency (Vmax/Km) allow the conversion of methanol, the product of methane activation, down to low concentrations.

“Ca. Methylomirabilis oxyfera” MDH contains PQQ as its prosthetic group. While the unambiguous presence of this cofactor in an XoxF-type large subunit was established only quite recently (43), the finding was expected. Except for nicotinoprotein methanol/alcohol dehydrogenase (3, 4), all of the MDHs known to date rely on PQQ as the catalytic center and PQQ-binding motifs have been identified before in XoxF MDHs (see Fig. S1A) (8, 32). However, the genome of “Ca. Methylomirabilis oxyfera” lacks known PQQ biosynthesis machinery. So, either the organism has evolved a novel biosynthetic pathway, which is unlikely considering the highly specific chemistry of the (oxygen-dependent) reactions involved (67), or the microorganism derives the cofactor from producers in the enrichment culture. Our finding that the supply of boiled cells of M. fumarariolicum strongly stimulated “Ca. Methylomirabilis oxyfera” culture activity and growth supports the latter view. Otherwise, the dependence on exogenous PQQ is not without precedent. For example, Escherichia coli and other enteric bacteria are incapable of PQQ synthesis, but the organisms readily incorporate the cofactor present in growth medium into their glucose dehydrogenase quinoproteins (68, 69). In addition, it has been known for quite some time that microorganisms that are capable of PQQ biosynthesis may excrete the compound, thereby stimulating the growth of other organisms (70). Still, it would be astonishing if “Ca. Methylomirabilis oxyfera,” with its unique life-style, depended on other microorganisms in the environment for the supply of an essential cofactor of one of its key enzymes.

The large subunit of purified MDH from “Ca. Methylomirabilis oxyfera” belongs to the XoxF MDHs. XoxF MDHs likely represent an ecologically highly relevant but grossly overlooked group of MDHs (8). Genes coding for these proteins are widely found in the genomes of methylotroph and methanotrophic bacteria and can be phylogenetically differentiated into at least five lineages (see Fig. S2 in the supplemental material) (8, 34). While hardly expressed under laboratory growth conditions (32), xoxF genes are among the most highly expressed genes in natural sys-
tems (35, 36). The reason for their long-time elusiveness is that XoxF apparently harbors lanthanide REEs, compounds that are rarely included in laboratory trace element solutions, at its catalytic site (40–43). The presence of a REE may confer on the proteins a catalytic efficiency that is superior to that of their commonly studied calcium-containing MxaFI MDH counterparts (8, 43). Typically, the XoxF MDHs isolated thus far lack a small subunit (39–43). In agreement with this, XoxF genes are not associated with genes encoding a small subunit (8). In these respects, MDH from “Ca. Methylomirabilis oxyf era,” being the first member of the XoxF MDH family to be documented, already might represent a variation on a still only emerging theme. First, enzyme preparations did not contain appreciable amounts of REEs. However, calcium was present, although part of it may have stemmed from the hydroxypatite used during purification. Nonetheless, the UV-visible light spectrum of “Ca. Methylomirabilis oxyf era” MDH, having an absorbance maximum at 445 nm (Fig. 3), is typical of a calcium-containing quinoprotein rather than a REE-containing MDH, in which the PQQ absorbance band is shifted to a longer wavelength (454 nm) (43). These considerations suggest that “Ca. Methylomirabilis oxyf era” MDH possesses calcium. A second striking difference between the “Ca. Methylomirabilis oxyf era” MDH and the XoxF MDHs known thus far is the presence in the former of a small subunit that is derived from the MxaFI m gene cluster (MDH-1) (Fig. 1). The function of the small subunit is not precisely known, but it has been suggested by us (8) that it enables MxaFI MDHs to properly coordinate calcium. Also in the XoxF MDH described here, the small subunit might have such a function. Obviously, the concerted action of different proteins from different gene clusters requires tight regulation. The mechanisms underlying regulation are a wide-open field of research, regarding not only “Ca. Methylomirabilis oxyf era” but also in other methylothrophs and methanotrophs that harbor multiple MxaFI and XoxF MDH systems in their genomes.

The presence of multiple MDH systems offers a methylothroph or methanotroph the means to adapt its metabolism in an optimal way to prevailing environmental conditions (availability of substrates and essential cofactors, like metals). The as yet poorly explored XoxF MDHs add to this by their (presumably) superior catalytic properties, yet utilizing difficult-to-access catalytic REEs (8). The hybrid heterotetrameric MDH described here provides “Ca. Methylomirabilis oxyf era” with a most efficient means to oxidize methanol in the pathway from methane to CO2.

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