

# Characterization of a Corrinoid Protein Involved in the C1 Metabolism of Strict Anaerobic Bacterium *Moorella thermoacetica*

Amaresh Das,<sup>1</sup> Zheng-Qing Fu,<sup>1</sup> Wolfram Tempel,<sup>1</sup> Zhi-Jie Liu,<sup>1,2\*</sup> Jessie Chang,<sup>1</sup> Lirong Chen,<sup>1</sup> Doowon Lee,<sup>1</sup> Weihong Zhou,<sup>1</sup> Hao Xu,<sup>1</sup> Neil Shaw,<sup>2</sup> John P. Rose,<sup>1</sup> Lars G. Ljungdahl,<sup>1</sup> and Bi-Cheng Wang<sup>1</sup>

<sup>1</sup>Department of Biochemistry and Molecular Biology, University of Georgia, Athens, Georgia 30602

<sup>2</sup>National Laboratory of Biomacromolecules, Institute of Biophysics, Chinese Academy of Sciences, Beijing 100101, China

**ABSTRACT** The strict anaerobic, thermophilic bacterium *Moorella thermoacetica* metabolizes C1 compounds for example CO<sub>2</sub>/H<sub>2</sub>, CO, formate, and methanol into acetate via the Wood/Ljungdahl pathway. Some of the key steps in this pathway include the metabolism of the C1 compounds into the methyl group of methylenetetrahydrofolate (MTHF) and the transfer of the methyl group from MTHF to the methyl group of acetyl-CoA catalyzed by methyltransferase, corrinoid protein and CO dehydrogenase/acetyl CoA synthase. Recently, we reported the crystallization of a 25 kDa methanol-induced corrinoid protein from *M. thermoacetica* (Zhou et al., Acta Crystallogr F 2005; 61:537–540). In this study we analyzed the crystal structure of the 25 kDa protein and provide genetic and biochemical evidences supporting its role in the methanol metabolism of *M. thermoacetica*. The 25 kDa protein was encoded by *orf1948* of contig 303 in the *M. thermoacetica* genome. It resembles similarity to MtaC the corrinoid protein of the methanol:CoM methyltransferase system of methane producing archaea. The latter enzyme system also contains two additional enzymes MtaA and MtaB. Homologs of MtaA and MtaB were found to be encoded by *orf2632* of contig 303 and *orf1949* of contig 309, respectively, in the *M. thermoacetica* genome. The *orf1948* and *orf1949* were co-transcribed from a single polycistronic operon. Metal analysis and spectroscopic data confirmed the presence of cobalt and the corrinoid in the purified 25 kDa protein. High resolution X-ray crystal structure of the purified 25 kDa protein revealed corrinoid as methylcobalamin with the imidazole of histidine as the  $\alpha$ -axial ligand replacing benzimidazole, suggesting base-off configuration for the corrinoid. Methanol significantly activated the expression of the 25 kDa protein. Cyanide and nitrate inhibited methanol metabolism and suppressed the level of the 25 kDa protein. The results suggest a role of the 25 kDa protein in the methanol metabolism of *M. thermoacetica*. Proteins 2007;67:167–176. © 2007 Wiley-Liss, Inc.

**Key words:** X-ray crystal structure; corrinoid protein; methanol metabolism

## INTRODUCTION

*Moorella thermoacetica* (formerly *Clostridium thermoaceticum*) is a thermophilic, anaerobic acetogenic bacterium, which using the autotrophic Wood/Ljungdahl acetyl-CoA pathway converts C1 compounds such as CO<sub>2</sub>, CO, formate, and methanol to acetate.<sup>1–4</sup> This involves a total synthesis of acetate from two moles of CO<sub>2</sub>. One mole of CO<sub>2</sub> is reduced via formate and tetrahydrofolate intermediates to 5-methyltetrahydrofolate, the methyl group of which is transferred onto the cobalt atom of a corrinoid iron sulfur protein (C/Fe-S) forming a Co-methylcorrinoid.<sup>5</sup> In the final step, the methylcorrinoid and coenzyme A (CoA) were condensed with another mole of CO<sub>2</sub> forming acetyl-CoA catalyzed by the bifunctional Ni-Ni enzyme carbon monoxide dehydrogenase/acetyl-CoA synthase (CODH/ACS).<sup>6–10</sup>

The first indication of the involvement of corrinoids in the synthesis of acetate by the acetyl-CoA pathway was obtained by Poston et al.,<sup>11</sup> who found that the methyl group of synthetically produced methylcobalamin was incorporated into the methyl group of acetate by cell extracts of *M. thermoacetica*. This led to an investigation of the corrinoid content of *M. thermoacetica*.<sup>12</sup> Out of 15 different corrinoids found, 11 were identified. The most abundant B<sub>12</sub>-derivatives were 5-methoxybenzimidazolylcobamide (Factor III<sub>m</sub>) and cobyric acid, which both were present mostly as their Co-5'-deoxyadenosyl (coenzyme) derivatives, and also as Co-methyl derivatives. When intact cells of *M. thermoacetica* were exposed to <sup>14</sup>CO<sub>2</sub> the Co-methyl groups of the two Co-methyl corri-

Grant sponsor: National Institutes of Health; Grant number: 1P50 GM62407; Grant sponsors: University of Georgia Research Foundation, Georgia Research Alliance; Grant sponsor: U.S. Department of Energy; Grant number: DE-FG02-93ER20127; Grant sponsor: Institute of Biophysics, Chinese Academy of Sciences, Start up research fund.

Wolfram Tempel's current address is Structural Genomics Consortium, University of Toronto, 100 College Street, Toronto, ON M5G 1L6, Canada.

\*Correspondence to: Zhi-Jie Liu, National Laboratory of Biomacromolecules, Institute of Biophysics, Chinese Academy of Sciences, 15 Datun Road, Beijing 100101, China. E-mail: zjliu@ibp.ac.cn

Received 23 December 2005; Revised 4 April 2006; Accepted 1 May 2006

Published online 8 January 2007 in Wiley InterScience (www.interscience.wiley.com). DOI: 10.1002/prot.21094

noids got labeled and their  $^{14}\text{C}$ -methyl groups were converted to the methyl group of acetate by extracts of *M. thermoacetica* fortified with pyruvate.<sup>13</sup>

It is now well established that derivatives of vitamin B<sub>12</sub> play important roles in bacterial C1 metabolism.<sup>14,15,7,16</sup> The cobalamin-dependent methionine synthase is the most studied methyltransferase (Mtr).<sup>17,18</sup> Other corrinoid-dependent transferases have been isolated from acetogens,<sup>19</sup> methanogens,<sup>20–24</sup> and methylo-trophs<sup>25,26</sup> and the methyl donors in these reactions include methyltetrahydrofolate, methyltetrahydromethanopterin, methanol, acetate, methylamines, methylethers, and halomethanes.

The corrinoid proteins isolated from acetogenic bacteria that participate in the Wood/Ljungdahl pathway of acetate biosynthesis have been reviewed.<sup>7</sup> Three corrinoid proteins have been isolated from *M. thermoacetica*. The best characterized corrinoid protein in the Wood/Ljungdahl pathway is the C/Fe-S, and as discussed above it transfers the methyl group of methyltetrahydrofolate to the CODH/ACS. It is a  $\alpha\beta$  dimer having two subunits of 33 and 55 kDa. The smaller subunit carries the corrinoid 5-methoxybenzimidazolylcobamide, whereas the larger subunit has the [4Fe-4S] cluster.<sup>5,19</sup> A second corrinoid protein designated MtvC isolated by Naidu and Ragsdale<sup>27</sup> is part of a three component vanillate O-demethylase system. This enzyme system may have a broad specificity and be involved in the transfer of methyl groups from a number of methoxylated aromatic compounds functioning as methyl donors.<sup>28</sup> A similar system has been described for the acetogen *Acetobacterium dehalogenans*.<sup>29,30</sup>

Recently we reported the preliminary crystallography study of a 25 kDa corrinoid protein.<sup>31</sup> Based on the N-terminal amino acid sequence of the protein it was identified to be a homolog of MtaC, which is a corrinoid protein and a component of the methanol:CoM Mtr system of methane producing archaea.<sup>20,32</sup> This enzyme system consists of three components MtaA, MtaB, and MtaC. MtaB catalyzes the transfer of the methyl group from methanol to the corrinoid cofactor of MtaC, while MtaA catalyzes the transfer of the methyl group from the corrinoid of MtaC to CoM. Genes encoding homologs of MtaA, MtaB, and MtaC are found to be present in the *M. thermoacetica* genome (<http://www.tigr.org>). Here we provide physiological, genetic, and structural evidence that the 25 kDa polypeptide is a corrinoid Mtr. The role of this protein in the methanol metabolism and acetate biosynthesis of *M. thermoacetica* has been discussed.

## MATERIALS AND METHODS

### Bacterial Strain and Growth Conditions

*Moorella thermoacetica* strain ATCC 39073 was grown on 200 mM methanol or 1% (wt/v) glucose as a carbon source at 58°C under 100% CO<sub>2</sub> gas in semi-defined Drake's minimal medium<sup>33</sup> in 125-mL bottles, 4-L flasks, or 100-L fermentors as previously described.<sup>34,35</sup> Cultures were harvested at mid to late log phase

(OD<sub>600</sub> ~ 1.0) by centrifugation at 6000g and stored at –80°C until used.

### Purification of the 25 kDa Protein

The 25 kDa corrinoid protein was purified from cytosolic extracts as described.<sup>31</sup>

### Assays and Measurements

UV-visible absorption spectra of fractions containing the corrinoid protein were recorded with a dual wavelength spectrophotometer (Shimadzu, model 2051PC). The sample preparations for spectral analysis were according to Ljungdahl et al.<sup>36</sup> Proteins were estimated using the Lowry method as described.<sup>34</sup> SDS-polyacrylamide gel electrophoresis of proteins was according to Laemmli.<sup>37</sup>

### DNA and RNA Sources, PCR, and Northern Hybridization

*M. thermoacetica* genomic DNA was isolated using Puregene DNA purification system (Gentra, Minneapolis, MN). Total RNA was isolated using RNeasy mini kit from Qiagen (Qiagen Valencia, CA). Prior to use RNA was treated with RNase-free DNase I (Roche Applied Sciences, Indianapolis, IN). For Dot-blot hybridization experiments RNA was denatured with formaldehyde (15% v/v in 5 × SSC at 60°C for 1 h) prior to application onto nylon membranes (ICN, Costa Mesa, CA). Hybridization (Northern or Dot-blot) experiments were carried out using the Genius system (Roche Applied Sciences) as described.<sup>38,39</sup> The PCR and the labeling of the PCR products with digoxigenin (DIG) were carried out using the FailSafe PCR System (EPICENTRE, Madison, WI). The DIG-labeled PCR product used as a probe in the hybridization experiments was a 462-bp fragment amplified from *orf1948* of contig 303 of the *M. thermoacetica* genome with 5'-TGACCAGGAGTTTGTGAGC-3' (forward primer) and 5'-CCGACGATTACTTTTACCCG-3' (reverse primer) using *M. thermoacetica* genomic DNA as template.

### N-Terminal Protein Sequence

To determine the N-terminal sequence of the 25 kDa protein the purified protein was subjected to SDS-PAGE and then trans-blotted onto PVDF membranes (Bio-Rad). After briefly stained with Coomassie Brilliant Blue the polypeptide was excised from the blot and sequenced at the Integrated Biotechnology Laboratories at the University of Georgia.

### Antibodies and Western Blotting Experiment

Polyclonal antibodies against the purified 25 kDa protein were raised in adult New Zealand white rabbit at the animal facility of the University of Georgia. Antibodies against CO dehydrogenase/acetyl-CoA synthase (CODH/ACS), Mtr, and corrinoid iron-sulfur protein (Co/Fe-S) were kindly provided by Steve Ragsdale of the

**TABLE I. Data Processing Statistics for the Refinement Data Set**

Resolution range (Å)	50.00–1.60 (1.66–1.60)
Wavelength (Å)	1.5798
Space group	P2 <sub>1</sub> 2 <sub>1</sub> 2
Cell dimensions (Å)	$a = 55.69, b = 62.74, c = 34.54$
Unique reflections	13939 (625)
Completeness (%)	84.1 (38.6)
$I/\sigma(I)$	46.8 (10.2)
$R_{\text{sym}}$ (%) <sup>a</sup>	7.7 (21.6)
Redundancy	5.7 (2.9)

$$^a R_{\text{sym}} = \sum_j |I_j - \langle I \rangle| / \sum_j \langle I \rangle.$$

University of Nebraska. Western blotting experiments were carried out according to Bio-Rad.

### Crystallization of the 25 kDa Protein and Collection of Diffraction Data

Crystallization and collection of a diffraction data to 1.9 Å resolution on a copper rotating anode source have been described previously.<sup>31</sup> The position of a single anomalous scatterer was determined by the program SHELXD in single wavelength anomalous scattering (SAS) mode.<sup>40</sup> Initial phases were calculated with the program SOLVE.<sup>41</sup> Phase improvement and automated model building were performed with RESOLVE.<sup>42</sup> A higher resolution data set was collected at a wavelength of 1.5798 Å at beamline 22ID of the Advanced Photon Source. A continuous sweep of 300 consecutive 1° oscillation images was recorded with a Mar300 CCD detector at a crystal-to-detector distance of 110 mm with an exposure of 4 s per image. Data reduction was carried out with the HKL2000 suite<sup>43</sup> (Table I). Further automated model building was carried out with ARP/wARP<sup>44,45</sup> using structure factor amplitudes<sup>46</sup> derived from the higher resolution data set with the program TRUNCATE of the CCP4 suite.<sup>47</sup> Iterative model validation, rebuilding, and refinement were carried out with MOLPROBITY,<sup>48,49</sup> XFIT<sup>50</sup> and the CCP4 program REFMAC5,<sup>51</sup> respectively. ARP/wARP and CCP4 programs were controlled through the CCP4I interface.<sup>52</sup> Coordinates of the refined model were deposited at the Protein Data Bank (PDB)<sup>53</sup> using the program PDB\_EXTRACT<sup>54</sup> with access code 1Y80.

### *M. thermoacetica* Genome Sequence

A draft annotated nucleotide sequence of the *M. thermoacetica* (ATCC39073) genome has been completed at the Joint Genome Institute Department of Energy and can be viewed at <http://www.tigr.org>.

## RESULTS

### Purification and Spectral Properties of the 25 kDa Protein

The 25 kDa corrinooid protein was purified from cytosolic extracts of methanol grown cells of *M. thermoacetica*

following ammonium sulfate precipitation, ion exchange chromatography, and gel filtration as described.<sup>31</sup> By size exclusion chromatography on Superose 12 column, an approximate molar mass of the protein was estimated to be 25 kDa (not shown). Since the predicted molar mass of the protein based on the amino acid sequence is 22,329 Da,<sup>31</sup> the purified protein as isolated is apparently a monomer in its native form. The amount of cobalt was estimated to be 0.9 mol per mol of the protein with negligible interferences from other metals. UV-visible spectra of the purified protein exhibited absorptions at 357 and 542 nm in oxidized form which were shifted to 363 and 551 nm following treatment with cyanide and to 369 and 580 nm after boiling with cyanide. These absorption maxima are typical for corrinooids as previously reported by Ljungdahl et al.<sup>36</sup>

### Identification of the 25 kDa Protein From the *M. thermoacetica* Genome Sequence

The N-terminal amino acid sequence of the purified 25-kDa protein was determined to be M(P)TYEELS-QAVFEGD. This sequence is identical to the predicted N-terminal amino acid sequence of the polypeptide encoded by *orf1948* of contig 303 of the *M. thermoacetica* genome. The *orf1948* is 630 bp long and encodes 210 amino acids (see Fig. 1). Analysis of the deduced amino acid sequence of *orf1948* revealed its similarity to MtaC, a component of the methanol:CoM Mtr system of methanogenic archaea,<sup>16</sup> and also to the C-terminal amino acid sequence of 5-methyltetrahydrofolate S-homocysteine Mtr from several bacteria including *Thermotoga maritima* (accession no. B72397, 35.1% identity), *Mycobacterium tuberculosis* (accession no. G70513, 34.1% identity), and *Bacillus halodurans* (accession no. B72397, 39.5% identity) (see Fig. 2). Both MtaC and the above enzymes shared a common sequence motif Asp-X-His-X-X-Gly-X<sub>41</sub>-Ser-X-Leu-X<sub>26-28</sub>-Gly-Gly, which has been rationalized as the signature for the corrinooids, and the His residue serves as the α-axial ligand to the corrinooid.<sup>15</sup> The above motif is also common to a subset of other B<sub>12</sub> enzymes including methyl-malonyl-CoA mutase, glutamate mutase, and methionine synthase.<sup>15</sup> In *M. thermoacetica* the methyl transfer reaction of the Wood/Ljungdahl pathway was catalyzed by methyltetrahydrofolate-dependent Mtr and the corrinooid iron-sulfur protein (C/Fe-S). The primary structure of C/Fe-S (ORF1921 of contig 303) lacks the above signature and has corrinooid with water instead of His as α-axial ligand. In the *M. thermoacetica* genome *orf1948* was surrounded by two additional ORFs, *orf1949* and *orf1947* (see Fig. 1). The *orf1949* is 1401 bp long encoding 476 amino acids, and *orf1947* is 801 bp long encoding 276 amino acids. The three ORFs were organized in the order *orf149*>*orf1948*>*orf1947* (see Fig. 1), which we now referred to as the corrinooid cluster of *M. thermoacetica*. The corrinooid cluster was located 3009 bp upstream of the CODH/ACS cluster (see Fig. 1) that contained the genes encoding CODH/ACS (ORFs 1919 and 1920), C/Fe-S (ORFs 1921 and 1924) and Mtr (ORF





Fig. 1. The organization of the genes of the corrinoid cluster and the relationship between the genes and the corresponding products derived from the annotated *M. thermoacetica* genome. The *acaA-accaF* cluster encoding CODH/ACS, C/Fe-S, and Mtr<sup>19</sup> was located 3907 bp upstream of the corrinoid cluster.

G1948	1	NPTY.EE...	.....	.....	.....	.....	LSQAVFEG	14
Mmaze_MtaC	1	MLKSKWRFN	MLDFTASLK	KVLTRYNVAL	EKANTPEDAA	EELYPKDELI	YPIAKAIFEG	60
Afulg_MtaC	1	MADVKE...	.....	.....	.....	.....	..FVQALADL	15
Mbark_MtaC	1	.....	MLDFTASLK	KVLTRYNVAL	EKALTPEEAA	EELYPKDELI	YPIAKAIFEG	50
F83853	621	SNLTLE.ER.	.....	.....	.....	.....	..LANYIVEG	636
B72397	561	PRAEVKEEK.	.....	.....	.....	.....	..LVEIILSG	577
G70513	641	G.LPLF.ER.	.....	.....	.....	.....	..LAQRIVDG	655
G1948	15	DEAQVVELTR	SLLSGGAEPL	EVINKGLIAG	HDRVGVLFEN	NEHFVPEVLH	SANAHNAGVE	74
Mmaze_MtaC	61	EEDDVIEGLE	AAIKAGKDFI	ALIDDALHVG	NGVVTIRLYDE	GIIFLPNVHM	SADAHLDGIE	120
Afulg_MtaC	16	DEAKTVELTK	KRVESGEDPF	TILEDV.RKA	TDIIGKRFE	GRYFVSDLM	AGEILKQVME	74
Mbark_MtaC	51	EEDDVVEGLQ	AAIEAGKDFI	DLIDDALHVG	NGVVIRLYDE	GVIFLPNVHM	SADAHLEGIE	110
F83853	637	SKDGLTEDLD	KALAKYDDPL	DIINGPLMNG	HDEVGRLFMN	NELIVAEVLQ	SAEVHKASVA	696
B72397	678	NRSELEKLVE	DFLKE.KDPL	SVIEEHLRPA	NERIGELYDK	GKIFLPQLIL	AAQTVKPVFD	736
G70513	656	ERNGLDADLD	EAMTQ.KPPL	QIINEHLLAG	NKTVGELFGS	GQMQLPFVLQ	SAEVHKAAVA	714
G1948	75	VVKQSQAQFD	HPS.VGKIVL	GTVEGDLHDI	GKNLVAMHLE	SGGFTVYNLG	VDIEPGKFVE	133
Mmaze_MtaC	121	FCKENSETAP	VT..KGTVVC	HVAEGDVHDI	GKNIVTALLR	ANGYNVVDLG	RDVPVDEVLN	178
Afulg_MtaC	75	ILRPLLGEKK	AES.KGKVVI	GTVEGDVHDI	GKNIVIALLE	AEGFEVVDIG	VDQPEAFVE	133
Mbark_MtaC	111	YCKENSGATP	KT..KGTVVC	HVAEGDVHDI	GKNIVTALLR	ANGYNVVDLG	RDVPAAEVL	168
F83853	697	HLEPHMEKKA	DDHGKGKIL	ATVKGDVHDI	GKNLVEIILS	NNGFRIVNLG	IKVTSNELIE	756
B72397	737	KLTSMLPSDS	QGETF...VI	ATVKGDVHDI	GKNIVASVIR	SSGYRVVDLG	KDVTSEIVE	793
G70513	715	YLEPHMERSD	DDSGKGRIVL	ATVKGDVHDI	GKNLVDIILS	NNGYEVVNI	IKQPIATILE	774
G1948	134	AVKKYQPDIV	GMSALLTTT	MNMKSTIDAL	IAAGLRDRVK	..VIVGGAPL	SQDFADEIGA	191
Mmaze_MtaC	179	AVANNPILV	TGTALHTTT	YAFKEVNDKL	LEKGYK..IP	..FACGGGAV	NQDFVSQYAL	234
Afulg_MtaC	134	AANQHNPVV	GLSGLLTAI	ESMKRTVEAL	RKAGYK..GK	..IIIGGGRT	SEEAKYTTGA	185
Mbark_MtaC	169	AVQKEKPIHL	TGTALHTTT	YAFKEVNDKL	LENGIK..IP	..FACGGGAV	NQDFVSQFAL	224
F83853	757	AVARENPDIAI	GLSGLLVKSA	QHMVLAQDL	KQQQIS..IP	..ILVGGGAL	TRKFTNTKIA	812
B72397	794	AVEKERPVAL	GLSAMHTTV	GRIKEVVEKL	KEKNLK..IP	..VIVGGASL	NEKLAKELGA	845
G70513	775	VAEDKSADVV	GMSGLLVKST	VVMKENLEEM	NTRGVA..EK	FPVLLGGGAL	TRSYVENDLA	832
G1948	192	DGYAPDAASA	TELCRQLLE.	.....	.....	.....	210	
Mmaze_MtaC	235	GVYGEAAADA	PKIADAIVAG	TTDIAALRDK	FHKH	.....	268	
Afulg_MtaC	190	DDWADDAVVG	VRIKALVGV	E.....	.....	.....	210	
Mbark_MtaC	225	GVYGEAAADA	PKIADAIAG	TTDVTREK	FHKH	.....	258	
F83853	813	PEYDGLVVYA	KDAMNGLELA	NKLNKPDE.	.....	.....	840	
B72397	850	DYAKNASEA	VKILKSLGR.	.....	.....	.....	868	
G70513	833	EIIQGEVH.	.....	.....	.....	.....	840	

Fig. 2. Multiple sequence alignment of the 25 kDa polypeptide (*orf1948*) from *M. thermoacetica* with MtaCs from *Methanosarcina mazei* Go 1 (accession no. NP\_633672); *Archaeoglobus fulgidus* DSM 4304 (accession no. NP\_068847); *Methanosarcina barkeri* (accession no. CAA69619), and 5-methyltetrahydrofolate S-homocysteine methyltransferases from *Bacillus halodurans* (strain C-125) (accession no. F83853); *Thermatoga maritima* (accession no. B72397); and *Mycobacterium tuberculosis* (accession no. G70513). Conserved residues were shown in bold face. Residues marked by asterisk belong to the conserved corrinoid-binding domain of the protein.

1925).<sup>19</sup> The polypeptide encoded by *orf1949* revealed similarity to MtaB (see Fig. 3), the second component of the methanol:CoM Mtr system, from several methanogenic archaea.<sup>20</sup> In methanogenic archaea, *mtaC* and *mtaB* were co-transcribed from a single polycistronic operon, while *mtaA* encoding the third component MtaA of the enzyme system was present separately at a distant location.<sup>20</sup> As described below *orf1949* and *orf1948* of *M. thermoacetica* were also co-transcribed from a polycistronic operon. Search for the gene encoding the third component of the Mtr system MtaA revealed its similarity to the polypeptide encoded by ORF2632 of contig 309 in the *M. thermoacetica* genome (see Fig. 4) as the deduced *orf2632* shows similarity to MtaAs from several

archaea. Therefore, homologs of all three components of the methanol:CoM Mtr system of methanogenic archaea are present in *M. thermoacetica*. The polypeptide encoded by *orf1947* of the corrinoid cluster revealed similarity (33.6% identical residues) to C/Fe-S of *M. thermoacetica*, and also to the N-terminal amino acid sequence of 5-methyltetrahydrofolate S-homocysteine Mtrs from several bacteria (not shown).

### Northern Blot Analysis and Regulatory Sequences

The *orf1949* and *orf1948* of the corrinoid cluster of *M. thermoacetica* were separated by 110 bp, while *orf1948*

**A**

G1949	1	MDYKPVKTF	ELEVKSLDDF	VYGIAPHPVK	AKNGHIVIGAG	TVYPEINMTL	PPMNIEESTM
Mmaze_MtaB	1	MAATR...FT	KMAYASADEM	TFGVSKYPVK	AGLGLLEIGAG	YTIPEVNYAP	RPEAG..ASK
Mbark_MtaB	1	MAAKR...YT	SMAYANADEM	TFGVSKYPVK	AGLDLEIGAG	YTIPEINYAP	RPEAG..ASK
Mace_MtaB	1	MAAKR...YT	SMAYASADEM	SFGVSKYPVK	AGLGLLEIGAG	YTIPEVNYAP	RPEAG..ASK
G1949	61	PEVRRQYAE	IEGILKRARD	LYAPGIIVEL	ELLPETTMKP	EWGIEINKIL	RDRMHEYEDK
Mmaze_MtaB	56	EKLIKEYERI	TTDIMARMVQ	VGFPVILET	EHVQOMSNNP	SWGAEVAHAQ	KTMEYEHDE
Mbark_MtaB	56	EKLIKEYERI	TTDVHMRMVQ	VGFPVILET	EHVQOMSNNP	SWGAEVAHAQ	KTMEYKHDE
Mace_MtaB	56	EKLKEYERI	TTDIMGRMVQ	VGFPVILET	EHVQOMSNNP	SWGAEVAHAQ	KTMEYEHDE
G1949	121	YGLKSLLRCT	PNDTREILRP	PLMKRGELLE	NMFITFEKCA	EDGADILSIE	STGGKEVHDE
Mmaze_MtaB	116	YGIKCALRHT	IGDIRENRDF	LQLRGDKYSV	FLEA.FEECA	KAGADLLSVE	SMGGKEVFEDY
Mbark_MtaB	116	YGIKCALRHT	IGDIRENREF	LQLRGDKYSV	FLEA.FEQCA	ENGADLLSVE	SMGGKEVFEDY
Mace_MtaB	116	YGIKCALRHT	IGDIRENRDF	LQLRGDKYSV	FLEA.FEECA	KSGADLLSVE	SMGGKEVFDH
G1949	181	ALVTCNIRKA	IFALGVLGVR	DMRFLWSNIV	RIAERTGAIA	GGDTACGFAN	TALALAEQGM
Mmaze_MtaB	175	AVLRNDIAGM	LYAIGCLGSI	DMELIWSDIS	AIAKKTGTVS	AGDTDCAQAN	TAMFIGGGLL
Mbark_MtaB	175	AVLRNDIPGL	LYSIGCLGSI	DMELIWTDIS	KIAKKTGTIS	AGDTDCAQAN	TAMFIGGGLL
Mace_MtaB	175	AVLRNDVAGM	LYAIGCLGSI	DMEMIWSDIA	AIAQKTGTVA	AGDTDCAQAN	TAMFIGGGLL
G1949	241	I PRVF...AA	VDRVATIPRS	LVAFEMGAIG	PKKDCGYEGP	YMKAIAGVPI	SMEGKTAACA
Mmaze_MtaB	235	DKNLAHTLAI	LARAIAPRS	LVAYECGAVG	PGKDCGYENV	VIKAITGMPM	TQEGKTSTCA
Mbark_MtaB	235	NKNLAHTIAV	IARAIAPRS	LVAYEAGAVG	PGKDCGYENI	IVKAITGMPM	TMEGKTSTCA
Mace_MtaB	235	DKNLAHTLAI	LARAIAPRS	LVAYECGAMG	PGKDCGYENI	IIKAITGKPM	TQEGKTSTCA
G1949	298	HLAIGNIAA	CVCDMWSNES	VQNVKLLSAP	APVVSTEQLI	YDCRIMNERA	ADGRSFALKM
Mmaze_MtaB	295	HSDVMGNLIM	QCCDCWSNES	VEYHGEFGGT	TVQCWGESLA	YDCALMNTAL	ETKNDKV..L
Mbark_MtaB	295	HSDVMGNLVM	QCCDCWSNES	VEYHGEFGGT	TVQCWSETLA	YDCALMNTAL	ETKNDKV..L
Mace_MtaB	295	HSDVLGNLIM	QCCDCWSNES	VEYHGEFGGT	TVQCWSETLA	YDCTIMNTAL	ETKNEKV..L

**B**

G1949	358	RDWLAASDSR	LDPQAYVLRP	DIVLEISQEL	VKEKDA.FIA	TKKAAALAAE	VIKRLARGE
Mmaze_MtaB	353	RDLLMLSDRY	RDPQAYVLAY	DNAYRIGQAI	VKDGDNIYLR	AKNAAIACCD	IVSEG.AAGK
Mbark_MtaB	353	RDLMLSDRY	RDPQAYMLAY	DNAYRVGQSI	VKDGDNIYLR	AKNAAIECCN	IIIEG.AAGK
Mace_MtaB	353	RDLFMLSDRY	RDPQGYVLAY	DNAYKVGEAI	VKDGEDITYLR	AKNAAVACCD	IVSEG.AAGK
G1949	417	VQVSSREKKW	LDIISSQIET	IPDDWEEFWY	EQKEL..DL	EKFRPEEYDL	EVIMARGASA
Mmaze_MtaB	412	LELSRFETKA	LADAKASLDS	LTDDMDKFMD	DCLTKYKSEV	KVFLPENYGF	.....
Mbark_MtaB	412	LELSRFETKA	LADAKAALEA	LPDDMDKFMD	DCLTKYKSEV	KVFKPENYGF	.....
Mace_MtaB	412	LELSRFETKA	LADAKASLDS	LTDDMDKFMD	DCLTKYKSEV	KVFLPENYGF	.....
G1949	475	GN					
Mmaze_MtaB		..					
Mbark_MtaB		..					
Mace_MtaB		..					

Fig. 3. Multiple sequence alignment of the protein encoded by *orf1949* of contig 309 of *M. thermoacetica* genome with MtaBs from *M. mazei* Go 1 (accession no. NP\_633671), *M. acetovorans* C2A (accession no. AAK07601) and *M. barkeri* (accession no. CAA69620). Conserved residues were shown in bold face.

and *orf1947* by 115 bp. A putative promoter sequence, 5'-ATGACC-N<sub>15</sub>-TTAAT-3', resembling the *E. coli* consensus  $\sigma$  promoter (5'-cTTGACA-N<sub>15-21</sub>-TATAaT-3')<sup>55</sup> was located 19-bp upstream of the *orf1949* start codon. No secondary promoter structure was apparent in the two intergenic regions, suggesting all three genes belong to a single polycistronic operon. To verify this presumption total RNA from *M. thermoacetica* was subjected to Northern hybridization with a 462-bp DIG-labeled DNA fragment amplified from *orf1948* as a probe. The Northern blots exhibit a strong hybridization signal at 3200 bp which is close to the combined size of *orf1947*, *orf1948*, and *orf1949* [Fig. 5(A)] including the intergenic regions 3009 bp, suggesting a polycistronic message for the corrino cluster. The hybridization signal was much stronger with RNA from methanol grown cells than with RNA from glucose-grown cells. Hybridization signals

were virtually undetectable with RNA from glucose plus nitrate grown cells grown which is expected since nitrate was reported to inhibit the C1 metabolism and the Wood/Ljungdahl pathway in *M. thermoacetica*.<sup>33</sup> RNA from cyanide-grown cells also failed to yield any hybridization (not shown) which could be due to inhibition of CO dehydrogenase/acetyl-CoA synthase,<sup>56</sup> the most crucial enzyme of the Wood/Ljungdahl pathway. The Northern blots show smear immediately following the hybridization signal at 3200 bp, indicating degradation of the transcripts, which is not unusual for polycistronic transcripts as reported earlier.<sup>35</sup> To verify the level of transcripts under different growth conditions, total RNA was subjected to dot-blot hybridization with the same probe used in Northern hybridization experiments. Results [Fig. 5(B)] show strong hybridization signals for RNA from methanol-grown cells than from glucose-grown or

G2632	9	TPKRRFLSAL	FGGRVDRTPV	ANPISLVTVE	IMERTGAYFP	DAHLDAEKMA	RLAATSYEVL
Mmaze_MtaA	8	TLKTRLLAAL	KGEFVDKVPV	CSVTQTGIVE	IMDVVGAPWP	EAHTNPEIMA	KLALANHEL
Mace_MtaA	8	TLKTRLLAAL	KGEFVDKVPV	CSVTQTGIVE	IMDEVGAPWP	EAHTNPEIMA	KLALANYELS
Mbark_MtaA	5	TPKERLYRAL	RKQGVDRMPA	VCFTQTATVE	QMEACGAYWP	EAHSDAERMA	TLAEAAHTVV
G2632	69	GFDTIMFVFS	AHTESAALGV	FVDWGDKMSW	PVNTSHFITD	PEQIVIPDSE	LEEPSMRT-V
Mmaze_MtaA	68	GLEAVRLPYC	LTVLVEAMGC	EINMGTKNRQ	PSVTGHFYPK	DLEGAAPVAD	LLQGRIPVW
Mace_MtaA	68	GLEAVRVPYC	LTVLVEAMGC	EINMGTKNRQ	PSVTGHFYPK	ALDGAAPVAD	LLQKGRIPAV
Mbark_MtaA	65	GFEAVRVFPD	ITAEAEFFGC	GKAGDLKQQ	PSVIKPSVKN	LEDLEKLKNY	NLKEGRF---
G2632	128	LDAIKILRSQ	YGDRVAIGK	TYGPWSLAYH	LVGTEFNLME	TILNPKARR	YLEVLLIASI
Mmaze_MtaA	128	LEAIKIIREK	VGPDVPIVGG	MEGPVTVASD	LVSVKSEMKW	SIKKTDLLEQ	ALDIATEASI
Mace_MtaA	128	LEAIKIIREK	VGPDVPIIGG	MEGPITVASD	LVSVKSEMKW	SIKSPDLFEQ	SLDLATAASI
Mbark_MtaA	122	-----	-----	-----	-----	-----	-----NV
G2632	188	LSAKAQIKAG	ADAILMGDH-	ATGDLVSAEY	YRDFLMKVHQ	YVTR---EVG	APIILHICGN
Mmaze_MtaA	188	IYANAMVEAG	ADVIAIADEV	ASPDLMSPDS	FRQFLKSRLLQ	KFAS---SVN	SVTVLHICGN
Mace_MtaA	188	AYANAMVEAG	ADIIAIADEV	ASPDLMSPDS	FKQYLQPRLLQ	KFSS---SVS	SVTVLHVCGN
Mbark_MtaA	124	AYAKAMVENG	ADTIAIIDPT	ASYELIGGEF	YEKALFYQK	KIVDAMKELD	VATVLHICGN
G2632	244	TTKFIPIYVE	AGFDAPHFDS	KVD-AKLAK	LAGNKMSLIG	NINNPTVLLA	GTPEVDKKE
Mmaze_MtaA	244	VNPILSDMAD	CGFEGLSVEE	KIGSAKKGKE	VIGTRARLVG	NVSSPTLLP	GPVDKIKAEA
Mace_MtaA	244	VNPILSYMAD	CGFEGLSVEE	KIGSVKKAKE	VIGTRARLIG	NISSPTLLP	GPVDKIKAES
Mbark_MtaA	184	TTNGLGINDK	TGVNGISVDQ	KVD-IKTATG	NVK-NALIVG	NLDPVAVLWN	GTPEEIAEVS
G2632	303	LYAIEAGVEI	VGPECAIPLT	TPLENILAIT	ETAKEYQIHK	KLGETQ	
Mmaze_MtaA	304	KEALEGGIDV	LAPGCGIAPM	TPLENVKALV	AARDEFYA--	-----	
Mace_MtaA	304	KQALADGVVD	LAPGCGIAPM	TPLENIKAMV	EARNEFYA--	-----	
Mbark_MtaA	242	KKVLDAGVGL	LTVGCGTVSM	TPTVNLQKMI	ECAKSHTY--	-----	

Fig. 4. Multiple sequence alignment of the protein encoded by *orf2632* of contig 309 of *M. thermoacetica* genome with MtaAs from *M. mazei* Go 1 (accession no. NP\_633094), *Methanosarcina acetovorans* C2A (accession no. NP\_619241), and *M. barkeri* (accession no. S62369). Conserved residues were shown in bold face.

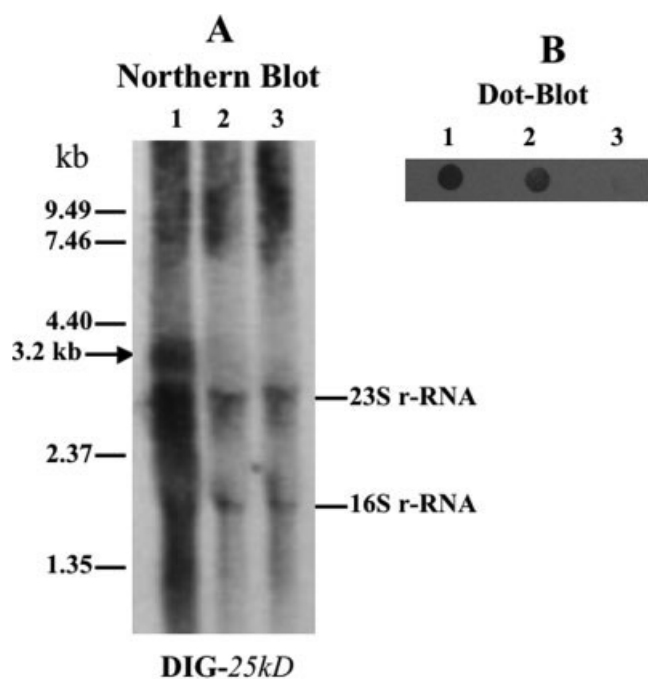


Fig. 5. Northern blot (A) and the dot blot (B) analysis of *M. thermoacetica* total RNA after hybridization with the DIG-labeled 462 bp PCR product amplified from *orf1948*. RNA was isolated from the bacterium grown on methanol (lane/dot 1), glucose (lane/dot 2) and glucose plus nitrate (lane/dot 3) as described in Materials and Methods.

glucose plus nitrate-grown cells. To check for any non-specific hybridization between the probe and the RNA, total RNA from these cells were also hybridized with DIG-labeled PCR-probe amplified from house-keeping gene *atpD* encoding the  $\beta$  subunit of  $F_1$ -ATPase.<sup>39</sup> Comparable hybridization signals were observed with RNA from methanol-, glucose-, and glucose plus nitrate-grown cells (not shown). These results suggest induction and strong activation of the corrino cluster by methanol.

### The Expression of the 25 kDa Protein Under Different Growth Conditions

Figure 6 shows higher level of expression of the 25 kDa protein in methanol-grown cells than in glucose-grown cells, and both nitrate and cyanide completely inhibited the expression of the protein. We compared the level of expression of the 25 kDa protein with that of other Wood/Ljungdahl pathway enzymes including the corrino protein C/Fe-S, and CODH and Mtr. Cyanide completely inhibited the expression of CODH, and significantly reduced the expression of C/Fe-S, Mtr, and the 25-kDa protein (see Fig. 7). In comparison to cyanide, nitrate had negligible effect on the expression of CODH, C/Fe-S, and Mtr in *M. thermoacetica*, also reported by Frostl et al.<sup>57</sup> The level of expression of the 25 kDa-protein was much higher in methanol-grown cells than in glucose-grown cells, which correlates well with the



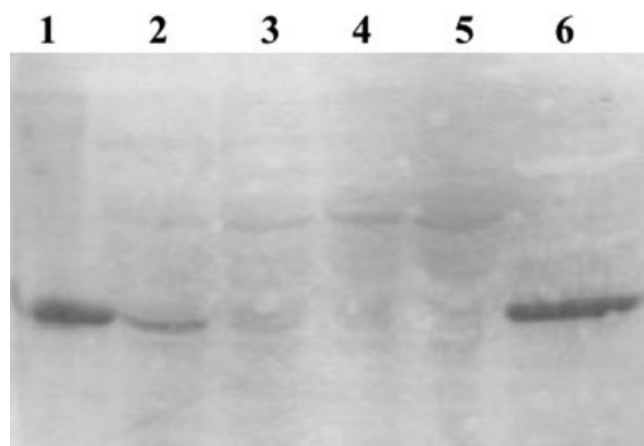


Fig. 6. Expression of the 25 kDa protein under different growth conditions. The purified 25 kDa protein (10  $\mu$ g, lane 6) and whole cell extracts (40  $\mu$ g per lane) of *M. thermoacetica* grown on methanol (lane 1), glucose (lane 2), glucose plus potassium cyanide (500  $\mu$ M, lane 3), glucose plus 5 mM  $\text{KNO}_3$  (lane 4), and glucose plus 15 mM  $\text{KNO}_3$  (lane 5) were subjected to SDS-PAGE, the proteins were transblotted from the SDS-gel onto PVDF membranes, and probed with antibodies against the 25 kDa protein by Western blotting experiments (details in the Materials and Methods).

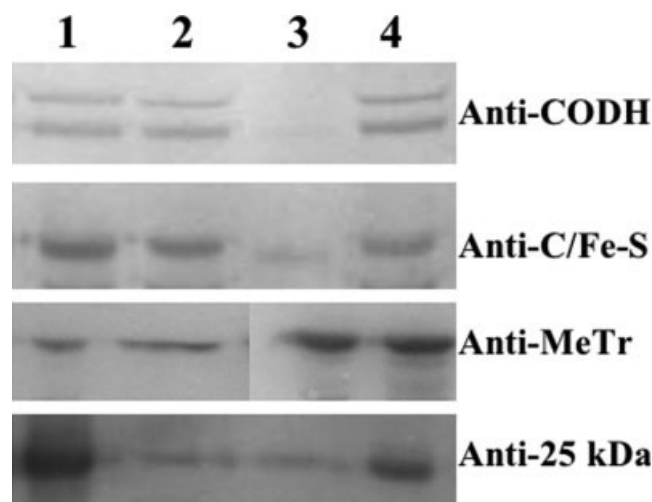


Fig. 7. Expression of the Wood/Ljungdahl pathway enzymes CODH/ACS (CO dehydrogenase/acetyl-CoA synthase), C/Fe-S (corrinoid iron-sulfur protein), and Mtr (methyltransferase) in *M. thermoacetica* grown on methanol (lane 1), glucose plus  $\text{KNO}_3$  (15 mM, lane 2), glucose plus KCN (500  $\mu$ M, lane 3) and glucose only (lane 4). Whole cell extracts (40  $\mu$ g per lane) were subjected to SDS-PAGE and the proteins were transblotted onto PVDF membranes as described in the legends to Fig. 8. Western blotting experiments were carried out with antibodies raised the 25 kDa protein, CODH/ACS, C/Fe-S, and Mtr as described in the Materials and Methods.

results of Northern hybridization experiments (see Fig. 5) suggesting a role of this protein in the methanol metabolism of *M. thermoacetica*. Having the similarities of the 25 kDa protein and those encoded by *orf1949* and *orf2632* with MtaC, MtaB, and MtaA, respectively, of the methanol:CoM Mtr methyltransferase system of methanogenic archaea, it is likely that *orf1948*, *orf1949*, and *orf2632* could function as a Mtr system in *M. thermoacetica*.

Winter-Ivey and Ljungdahl<sup>58</sup> suggested that the synthesis of acetyl-CoA from methanol could occur via direct transfer of the methyl group of methanol to CODH/acetyl-CoA synthase. However, this interpretation of the results needs verification at the enzyme level.

### Analysis of the Crystallographic Data and the Structural Configuration of the 25 kDa Protein

The cobalt ion of the corrinoid facilitated protein structure determination by acting as an anomalous scatterer during diffraction. The preparation of a heavy atom derivative was therefore not required. Using a data set obtained from a copper X-ray source, the cobalt ion could indeed be located and initial phases calculated using the SAS method.<sup>59</sup> Crystals contained an N-terminal truncation<sup>31</sup> of the full-length protein and residues 85–209 were located in the electron density map (see Fig. 8). The model was refined using data to 1.7 Å resolution and showed both good fit to the experimental data and reasonable geometry (Table II). At 30% sequence identity over the alignment region, the overall structure aligns with residues 743–871 of PDB entry 1BMT,<sup>60</sup> the model of methionine synthetase from *E. coli*, with an RMSD of 1.1 Å.<sup>61</sup> The location of the corrinoid is outlined in good detail by strong electron density (see Fig. 9). Differentiation from cobalamin is unambiguous due to strong density for the 5-methoxy group and the lack of density in the 4-position of the benzimidazole moiety of the corrinoid protein. The benzimidazole moiety is displaced from the cobalt ion by the imidazole moiety of residue His-101 (see Fig. 9) resulting in the “base-off” configuration of the corrinoid. Additional polar interactions between the corrinoid and the protein have been identified and are listed in Table III including the interactions of some of the amino acid residues that belong to the so-called the corrinoid signature motif of the polypeptide.

### DISCUSSION

Based on the primary structure the 25 kDa protein appears to be a homolog of MtaC, the corrinoid of the methanol:CoM Mtr system of methane-producing archaea.<sup>20</sup> The latter enzyme system consists of two additional enzymes MtaB and MtaA. In the *M. thermoacetica* genome homologs of MtaB and MtaA were found to be encoded by *orf1949* of contig 303 and *orf2632* of contig 309, respectively. In methanogenic archaea *mtaC* and *mtaB* are present in a polycistronic operon while *mtaA* in a separate loci. A similar organization is also found for the corresponding genes in *M. thermoacetica* (see Fig. 1). The 25 kDa polypeptide has no homology to either Mtr or the corrinoid iron-sulfur protein C/Fe-S of the Wood/Ljungdahl pathway which catalyzes methyl transfer reactions from methyltetrahydrofolate to synthesize acetyl-CoA.<sup>5</sup> The expression of the 25 kDa protein was induced and activated by methanol (see Fig. 6) suggesting a role of this protein in the methanol metabolism of *M. thermoacetica*. In methanogenic archaea, methane

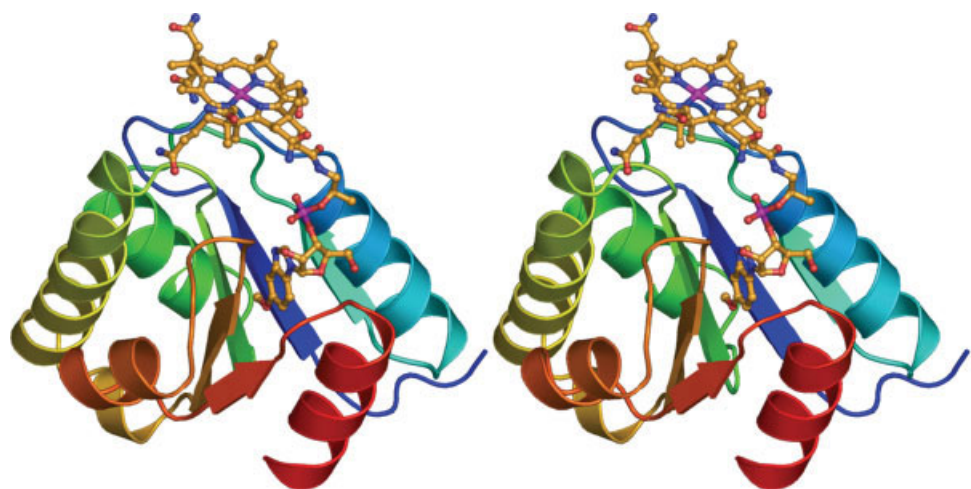


Fig. 8. Stereo view of the crystal structure of the 25 kDa corrinoid protein from *M. thermoacetica*. The image of the crystals was generated with software program PYMOL (DeLano, 2002).

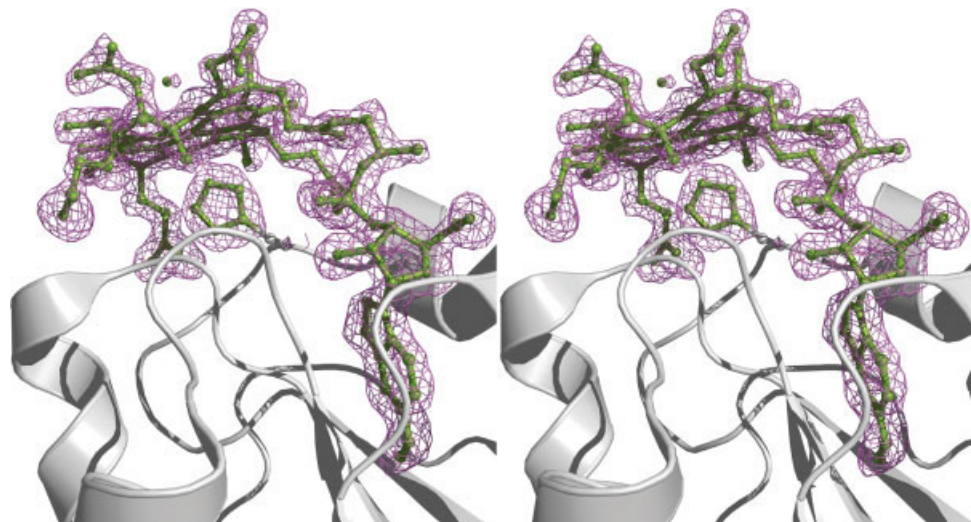


Fig. 9. Coordination of the corrinoid-bound cobalt ion with the ligand of the 25 kDa protein. The corrinoid is seen in the “base-off” configuration where Ne2 of His-101 displacing the benzimidazol moiety of the corrinoid as the ligand. Atoms shown in green were omitted in the calculation of a model-phased Fo-Fc electron density map shown here in magenta contoured at 2σ around the omitted atoms.

TABLE II. Statistics for the Protein Model

Resolution range (Å)	41.63–1.70
Total number of HKLs used (free)	12184 (528)
$R_{\text{work}}$ ( $R_{\text{free}}$ ) <sup>a</sup>	17.4% (21.0%)
Mean/Wilson B factor (Å <sup>2</sup> )	14.1/14.0
Total number of refined atoms (water)	1081 (72)
RMSD from ideal bond lengths (Å)	0.015
RMSD from ideal bond angles (°)	2.0

<sup>a</sup> $R_{\text{work}} = \Sigma ||F_{\text{obs(work set)}}| - k|F_{\text{cal}}|| / \Sigma |F_{\text{obs(work set)}}|$ ;  $R_{\text{free}} = \Sigma ||F_{\text{obs(test set)}}| - k|F_{\text{cal}}|| / \Sigma |F_{\text{obs(test set)}}|$ .

TABLE III. Bond Distances for Polar Protein–(solvent)–Corrinoid Interactions

Residue			Residue			Distance (Å)
Type	ID	Atom	Type	ID	Atom	
Gly	104	N	Wat	401	O	2.90
Wat	401	O	B1M	301	O4	2.61
B1M	301	O7R	Ala	179	N	3.13
Ser	146	OG	B1M	301	N3B	2.87
B1M	301	CO	His	101	NE2	2.42
B1M	301	CO	Wat	458	O	2.66

is produced from methanol catalyzed by MtaA, MtaB and MtaC.<sup>20</sup> Since acetate, not methane, is the final product of methanol metabolism in *M. thermoacetica* (Das and vanHoek, unpublished) it is likely that the

methyl transfer reactions from methanol catalyzed by the proteins encoded by the corrinoid cluster could be coupled to acetate biosynthesis. The 25 kDa protein was poorly expressed under nonacetogenic conditions for



example in the presence of nitrate or cyanide (Figs. 6 and 7) which block acetogenesis in *M. thermoacetica*.<sup>56,57</sup> On the other hand, CODH, C/Fe-S and Mtr were all expressed in the presence of nitrate but not in the presence of cyanide (see Fig. 7). These results suggest a specific role of the 25 kDa protein in acetate biosynthesis.

Naidu and Ragsdale<sup>27</sup> reported an inducible three component aromatic *O*-demethylase system similar to that of methanol:CoM Mtr of methanogenic archaea in *M. thermoacetica*.<sup>20</sup> The three components of the *M. thermoacetica* *O*-demethylase system designated MtvA, MtvB and MtvC, were shown to catalyze direct transfer of the *O*-methyl group from methoxylated aromatic compounds for example syringate to the one carbon carrier tetrahydrofolate. The corrinoid protein of the aromatic *O*-demethylase from *M. thermoacetica* was shown to be MtvC. It was shown that MtvB catalyzed the transfer of the methyl group from phenylmethyl ether to the cobalt center of MtvC, and MtvA catalyzed the transfer of the methyl group from MtvC to tetrahydrofolate forming methyltetrahydrofolate. Methyltetrahydrofolate then served as the methyl donor in the synthesis of acetyl-CoA from CO and CoA catalyzed by Mtr, C/Fe-S, and CODH/ACS.<sup>8</sup> The N-terminal sequence of MtvC was reported to be MLTDTL(S)KAMAELEEEQ(V)LA which did not match the N-terminal sequence of the 25 kDa protein but matched exactly with the N-terminal sequence of the polypeptide encoded by *orf223* of contig270 of the *M. thermoacetica* genome. The *orf223* is preceded by two additional ORFs, *orf221* and *orf222*, neither of which has similarity to *mtaA* and *mtaB* of methanogenic archaea or *orf1949* and *orf2632* of *M. thermoacetica*. The predicted amino acid sequence of MtvC deduced from *orf223* shared 33% identical residues with that of the 25 kDa protein including the highly conserved corrinoid-binding motif Asp-X-His-X-X-Gly-X<sub>41</sub>-Ser-X-Leu-X<sub>26-28</sub>-Gly-Gly (not shown). Based on these similarities it is assumed that the functions of *orf1949*, *orf1948*, and *orf2632* could mimic that of the aromatic *O*-demethylase system of *M. thermoacetica*.<sup>20</sup> Therefore, ORF1949 (equivalent to MtvB) could catalyze the transfer of the methyl group from methanol to the 25 kDa protein (equivalent to MtvC) while ORF2632 (equivalent to MtvA) catalyzes the transfer of the methyl group from 25 kDa protein to tetrahydrofolate forming methyltetrahydrofolate. Finally acetyl-CoA could be formed from methyltetrahydrofolate following condensation with CO and CoA catalyzed by Mtr, C/Fe-S, and CODH/ACS as previously described.<sup>8</sup> All the enzymes, essential for the synthesis of acetyl-CoA from methyltetrahydrofolate including CODH/ACS, C/Fe-S, and Mtr were expressed in methanol-grown cells (see Fig. 7), suggesting their involvement in the acetate biosynthesis from methanol. Analysis of the *M. thermoacetica* genome revealed several genes encoding putative corrinoid proteins, the function of which could be coupled methyl transfer reactions from a variety of naturally occurring compounds. This metabolic potential of *M. thermoacetica* and other

acetogens led them to inhabit virtually any anoxic environment in nature.<sup>1</sup>

## ACKNOWLEDGMENTS

Crystallographic data were collected at Southeast Regional Collaborative Access Team (SER-CAT) 22-ID beamline at the Advanced Photon Source, Argonne National Laboratory. Supporting institutions may be found at [www.ser-cat.org/members.html](http://www.ser-cat.org/members.html). Use of the Advanced Photon Source was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, under Contract No. W-31-109-Eng-38.

## REFERENCES

1. Drake HL. Acetogenesis, acetogenic bacteria, and the acetyl-CoA "Wood/Ljungdahl pathway": past and current perspectives. In: Drake HL, editor. *Acetogenesis*. New York, NY: Chapman & Hall; 1994. pp 3–60.
2. Drake HL, Daniel SL. Physiology of the thermophilic acetogen *Moorella thermoacetica*. *Res Microbiol* 2004;155:869–883.
3. Ljungdahl LG. The autotrophic pathway of acetate synthesis in acetogenic bacteria. *Annu Rev Microbiol* 1986;40:415–450.
4. Wood HG, Ljungdahl LG. Autotrophic character of the acetogenic bacteria. In: Shively JM, Barton LL, editors. *Variations in autotrophic life*. San Diego, CA: Academic Press; 1991. pp 201–250.
5. Ragsdale SW, Lindahl PA, Munck E, Massbauer, EPR, and optical studies of the corrinoid/iron-sulfur protein involved in the synthesis of acetyl coenzyme A by *Clostridium thermoaceticum*. *J Biol Chem* 1987;262:14289–14297.
6. Seravalli J, Xiao Y, Gu W, Cramer SP, Antholine WE, Krymov V, Gerfen GJ, Ragsdale SW. Evidence that NiNi Acetyl-Co A Synthase is active and that the CuNi enzyme is not. *Biochemistry* 2004;43:3944–3955.
7. Ragsdale SW. The acetogenic corrinoid proteins. In: Banerjee R, editor. *Chemistry and biochemistry of B<sub>12</sub>*. Vol. 1. New York: Wiley; 1999. pp 633–695.
8. Ragsdale SW, Kumar M. Nickel-containing carbon monoxide dehydrogenase/acetyl-CoA synthase. *Chem Rev* 1996;96:2515–2540.
9. Lenhert PG, Hodgkin DC. Structure of the 5,6-dimethyl-benzimidazolylcobamide coenzyme. *Nature* 1961;192:937–938.
10. Doukov TI, Iverson TM, Seravalli J, Ragsdale SW, Drennan CL. A Ni-Fe-Cu center in a bifunctional carbon monoxide dehydrogenase/acetyl-Co A synthase. *Science* 2002;298:567–572.
11. Poston JM, Kuratomi K, Stadtman ER. Methyl-vitamin B<sub>12</sub> as a source of methyl groups for the synthesis of acetate by cell free extracts of *Clostridium thermoaceticum*. *Ann NY Acad Sci* 1964; 112:804–806.
12. Irion E, Ljungdahl L. Isolation of Factor III<sub>m</sub> coenzyme and cobyrinic acid coenzyme plus other B<sub>12</sub> factors from *Clostridium thermoaceticum*. *Biochemistry* 1965;4:2780–2790.
13. Ljungdahl L, Irion E, Wood HG. Total synthesis of acetate from CO<sub>2</sub>. I. *co*-Methylcobyrinic acid and *co*-(methyl)-5-methoxybenzimidazolylcobamide as intermediates with *Clostridium thermoaceticum*. *Biochemistry* 1965;4:2771–2780.
14. Krautler B. Chemistry of methylcorrinoids related to their roles in bacterial C1 metabolism. *FEMS Microbiol Lett* 1990;87:349–354.
15. Ludwig ML, Matthews RG. Structure-based perspectives on B<sub>12</sub>-dependent enzymes. *Annu Rev Biochem* 1997;66:269–313.
16. Banerjee R, Ragsdale SW. The many faces of vitamin B<sub>12</sub>: catalysis by cobalamin-dependent enzymes. *Annu Rev Biochem* 2003; 72:209–247.
17. Goulding CW, Postigo D, Matthews RG. Cobalamin-dependent methionine synthase is a modular protein with distinct regions for binding homocysteine, methyltetrahydrofolate, cobalamin, and adenosylmethionine. *Biochemistry* 1997;35:8082–8091.
18. Matthews RC. Cobalamin-dependent methyltransferases. *Acc Chem Rev* 2001;34:681–689.
19. Lu WP, Schiau I, Cunningham JR, Ragsdale SW. Sequence and expression of the gene encoding the corrinoid/iron-sulfur protein from *Clostridium thermoaceticum* and reconstitution of the recombinant protein to full activity. *J Biol Chem* 1993;268:5605–5614.

20. Ding Y-H, Zhang SP, Tomb JF, Ferry JG. Genomic and proteomic analyses reveal multiple homologs of genes encoding enzymes of the methanol:coenzyme M methyltransferase system that are differentially expressed in methanol- and acetate-grown *Methanosarcina thermophila*. *FEMS Microbiol Lett* 2002; 215:127–132.
21. Kremer JD, Cao X, Krzycki J. Isolation of two novel corrinoid proteins from acetate-grown *Methanosarcina barkeri*. *J Bacteriol* 1993;175:4824–4833.
22. Kruer M, Haumann M, Meyer-Klaucke W, Thauer RK, Dau H. The role of zinc in the methylation of the coenzyme M thiol group in methanol:coenzyme M methyltransferase from *Methanosarcina barkeri*. *Eur J Biochem* 2002;269:2117–2223.
23. Sauer KR, Thauer K. Methanol:coenzyme M methyltransferase from *Methanosarcina barkeri*: identification of the active-site histidine in the corrinoid-harboring subunit MtaC by site-directed mutagenesis. *Eur J Biochem* 1998;253:698–705.
24. van der Meijden P, te Brommelstroet BW, Poirot CM, van der Drift C, Vogels GD. Purification and properties of methanol:5-hydroxybenzimidazolylcobamide methyltransferase from *Methanosarcina barkeri*. *J Bacteriol* 1984;160:629–635.
25. Coulter C, Hamilton JTG, McRoberts WC, Kulakov L, Larkin MJ, Harper DB. Halomethane:bisulfide/halide ion methyltransferase, an unusual corrinoid enzyme of environmental significance isolated from an aerobic methylotroph using chloromethane as the sole carbon source. *Appl Environ Microbiol* 1999; 65:4301–4312.
26. Studer A, Vuilleumie S, Leisinger T. Properties of the methylcobalamin: H<sub>4</sub>folate methyltransferase involved in chloromethane utilization by *Methylobacterium* sp. strain CM4. *Eur J Biochem* 1999;264:242–249.
27. Naidu D, Ragsdale SW. Characterization of a three-component vanillate O-demethylase from *Moorella thermoacetica*. *J Bactiol* 2001;183:3276–3281.
28. Daniel SL, Keith ES, Yang H, Lin Y-S, Drake HL. Utilization of methoxylated aromatic compounds by the acetogen *Clostridium thermoaceticum*: expression and specificity of the CO-dependent O methylating activity. *Biochem Biophys Res Commun* 1991; 180:416–422.
29. Engelmann T, Kaufman F, Diekert G. Isolation and characterization of a veratrol:corrinoid protein methyl transferase from *Acetobacterium dehalogenans*. *Arch Microbiol* 2001;175:376–383.
30. Kaufmann F, Wohlfarth G, Diekert G. O-demethylase from *Acetobacterium dehalogenans*. Substrate specificity and function of the participating proteins. *Eur J Biochem* 1998;253:706–711.
31. Zhou W, Das A, Liu Z-J, Chang J, Chen L, Lee D, Tempel W, Rose JP, Ljungdahl LG, Wang B-C. Isolation, crystallization and preliminary X-ray analysis of a methanol-induced corrinoid protein from *Moorella thermoacetica*. *Acta Crystallogr Sect F* 2005; 61:537–540.
32. van der Meijden P, Heythuysen HJ, Pouwels A, Houwen F, van der Drift C, Vogels GD. Methyltransferases involved in methanol conversion by *Methanosarcina barkeri*. *Arch Microbiol* 1983; 134:238–242.
33. Seifritz C, Daniel SL, Gossner A, Drake HL. Nitrate as a preferred electron sink for the acetogen *Clostridium thermoaceticum*. *J Bacteriol* 1993;175:8008–8013.
34. Das A, Ivey DM, Ljungdahl LG. Purification and reconstitution into proteoliposomes of the F<sub>1</sub>F<sub>0</sub> ATP synthase from the obligately anaerobic gram positive bacterium *Clostridium thermoautotrophicum*. *J Bacteriol* 1997;179:1714–1720.
35. Das A, Silaghi-Dumitrescu R, Ljungdahl LG, Kurtz DM, Jr. Cytochrome bd oxidase and dioxygen tolerance of the strictly anaerobic bacterium, *Moorella thermoacetica*. *J Bacteriol* 2004;187: 2020–2029.
36. Ljungdahl LG, LeGall J, Lee J-P. Isolation of a protein containing tightly bound 5-methoxybenzimidazolylcobamide (factor III<sub>m</sub>) from *Clostridium thermoaceticum*. *Biochemistry* 1973;12:1802–1808.
37. Laemmli UK. Cleavage of structural proteins during the assembly of the head of bacteriophage T4. *Nature* 1970;227:680–685.
38. Das A, Ljungdahl LG. *Clostridium pasteurianum* F<sub>1</sub>F<sub>0</sub> ATP synthase: genetic composition, primary structure and some unusual properties. *J Bacteriol* 2003;185:5527–5535.
39. Das A, Ljungdahl LG. Composition and primary structure of the F<sub>1</sub>F<sub>0</sub> ATP synthase from the obligately anaerobic bacterium *Clostridium thermoaceticum*. *J Bacteriol* 1997;179:3746–3755.
40. Schneider TR, Sheldrick GM. Substructure solution with SHELXD. *Acta Cryst D* 2002;58:1772–1779.
41. Terwilliger TC, Berendzen J. Automated MAD and MIR structure solution. *Acta Cryst D* 1999;55:849–861.
42. Terwilliger TC. Automated structure solution, density modification and model building. *Acta Crystallogr D Biol Crystallogr* 2002;58:1937–1940.
43. Otwinowski Z, Minor W. Processing of X-ray diffraction data collected in oscillation mode. *Methods Enzymol* 1997;276:307–326.
44. Perrakis A, Morris R, Lamzin VS. Automated protein model building combined with iterative structure refinement. *Nat Struct Biol* 1999;6:458–463.
45. Perrakis A, Harkiolaki M, Wilson KS, Lamzin VS. ARP/wARP and molecular replacement. *Acta Cryst D* 2001;57:1445–1450.
46. French S, Wilson K. On the treatment of negative intensity observations. *Acta Cryst A* 1978;34:517–525.
47. Winn MD. An overview of the CCP4 project in protein crystallography: an example of a collaborative project. *J Synchrotron Radiat* 2003;10:23–25.
48. Davis IW, Murray LW, Richardson JS, Richardson DC. MOL-PROBITY: structure validation and all-atom contact analysis for nucleic acids and their complexes. *Nucleic Acids Res* 2004;32: W615–W619.
49. Lovell SC, Davis IW, Arendall GW, de Bakker PI, Word JM, Prisant MG, Richardson JS, Richardson DC. Structure validation by C $\alpha$  geometry:  $\phi$ ,  $\psi$  and C $\beta$  deviation. *Proteins* 2003;50:437–450.
50. McRee DE. XtalView/Xfit—A versatile program for manipulating atomic coordinates and electron density. *J Struct Biol* 1999;125: 156–165.
51. Murshudov GN, Vagin AA, Dodson EJ. Refinement of macromolecular structures by the maximum-likelihood method. *Acta Cryst D* 1997;53:240–255.
52. Potterton E, Briggs P, Turkenburg M, Dodson E. A graphical user interface to the CCP4 program suite. *Acta Cryst D* 2003; 59:1131–1137.
53. Berman HM, Westbrook J, Feng Z, Gilliland G, Bhat TN, Weissig H, Shindyalov IN, Bourne PE. The Protein Data Bank. *Nucleic Acids Res* 2000;28:235–242.
54. Yang H, Guranovic V, Dutta S, Feng Z, Berman HM, Westbrook JD. Automated and accurate deposition of structures solved by X-ray diffraction to the Protein Data Bank. *Acta Cryst D* 2004; 60:1833–1839.
55. Hawley DK, McClure WR. Compilation and analysis of *Escherichia coli* promoter DNA sequences. *Nucleic Acids Res* 1983;25:2237–2255.
56. Anderson ME, Lindahl PA. Organization of clusters and internal electron pathways in CO dehydrogenase from *Clostridium thermoaceticum*: relevance to the mechanism of catalysis and cyanide inhibition. *Biochemistry* 1994;26:8702–8711.
57. Frostl JM, Seifritz C, Drake HL. Effect of nitrate on the autotrophic metabolism of the acetogens *Clostridium thermoautotrophicum* and *Clostridium thermoaceticum*. *J Bacteriol* 1996;178: 4597–4603.
58. Winters DK, Ljungdahl LG. PQQ-dependent methanol dehydrogenase from *Clostridium thermoautotrophicum*. In: Jongejan JA, Duine JA, editors. PQQ and quinoproteins. Boston: Kluwer; 1989. pp 35–39.
59. Wang BC. Resolution of phase ambiguity in macromolecular crystallography. *Methods Enzymol* 1985;115:90–112.
60. Drennan CL, Huang S, Drummond JT, Matthews RG, Lidwig M. How a protein binds B12: a 3.0 Å X-ray structure of B12-binding domains of methionine synthase. *Science* 1994;266: 1669–1674.
61. Holm L, Sander C. Protein structure comparison by alignment of distance matrices. *J Mol Biol* 1993;233:123–138.