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Purified particulate methane monooxygenase from *Methylococcus capsulatus* (Bath) is a dimer with both mononuclear copper and a coppercontaining cluster

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Abstract

Particulate methane monooxygenase (pMMO) is a membrane-bound enzyme that catalyzes the oxidation of methane to methanol in methanotropic bacteria. Understanding how this enzyme hydroxylates methane at ambient temperature and pressure is of fundamental chemical and potential commercial importance. Difficulties in solubilizing and purifying active pMMO have led to conflicting reports regarding its biochemical and biophysical properties, however. We have purified pMMO from *Methylococcus capsulatus* (Bath) and detected activity. The purified enzyme has a molecular mass of ~200 kDa, probably corresponding to an $\alpha_2\beta_2\gamma_2$ polypeptide arrangement. Each 200 kDa pMMO complex contains 4.8 ± 0.8 copper ions and 1.5 ± 0.7 iron ions. Electron paramagnetic resonance (EPR) spectroscopic parameters corresponding to 40-60% of the total copper are consistent with the presence of a mononuclear type 2 copper site. X-ray absorption near edge (XANES) spectra indicate that purified pMMO is a mixture of Cu(I) and Cu(II) oxidation states. Finally, extended X-ray absorption fine structure (EXAFS) data are best fit with oxygen/nitrogen ligands and a 2.57 Å Cu-Cu interaction, providing the first direct evidence for a copper-containing cluster in pMMO. Abbreviations: pMMO, particulate methane monooxygenase; EPR, electron paramagnetic resonance; 12DM, dodecyl-β-D-maltoside; EXAFS, extended X-ray absorption fine structure.

Introduction

One of the great challenges to the chemical and engineering communities is the selective oxidation of methane to methanol. Although world reserves of petroleum and natural gas are comparable, methane is used far less efficiently as an energy source because it has a low energy density and is hazardous and expensive to transport (1). Conversion of methane to a liquid such as methanol would solve this problem, but current industrial processes for this transformation are costly and inefficient, requiring high temperatures and pressures (2, 3). By contrast, methanotrophic bacteria (4) oxidize methane to methanol at ambient temperature and pressure using methane monooxygenase (MMO) enzyme systems (5). Only one other enzyme, ammonia monooxygenase (6), can activate the C-H bond in methane (104 kcal/mole). Cytochromes P-450 (7) and copper monooxygenases (8) oxidize larger, more reactive hydrocarbons, but cannot hydroxylate methane. Therefore, a detailed understanding of biological methane oxidation is of both fundamental chemical and potential commercial importance.

All methanotrophs produce a membrane-bound methane monooxygenase called particulate methane monooxygenase (pMMO) (5). Under conditions of low copper availability, several strains also express a soluble enzyme (sMMO) (9), which contains a catalytic diiron center (10). Whereas the structure, biochemistry, and mechanism of sMMO are well understood (11), studies of pMMO are less advanced due to difficulties in solubilizing and purifying active enzyme. The *Methylococcus capsulatus* (Bath) pMMO comprises three polypeptides, the α (~47 kDa), β (~24 kDa), and γ (~22 kDa) subunits, encoded by the *pmoB*, *pmoA*, and *pmoC* genes, respectively (12, 13). It is not known how these three polypeptides are arranged in the pMMO holoenzyme. According to radiolabeling experiments with the suicide substrate acetylene, the active site is located on the β subunit (14, 15) and may also involve the α subunit (14).

The metal composition of the pMMO active site is the subject of much controversy. For purified samples, stoichiometries ranging from 2 (16) to 15 (14, 17, 18) copper ions per $\alpha\beta\gamma$ complex have

been reported. Based on electron paramagnetic resonance (EPR) spectroscopic studies of membranebound (19, 20) and purified (17) *M. capsulatus* (Bath) pMMO, Chan and co-workers propose that the pMMO copper ions are organized in multiple trinuclear clusters composed of one Cu(II) and two Cu(I) ions. These sites are hypothesized to fall into two functional classes, catalytic and electron transfer (20). The primary evidence for these trinuclear clusters derives from the analysis of the hyperfine splitting pattern of an isotropic signal at g=2.06 (19). A similar signal has been detected for *Methylosinus trichosporium* OB3b pMMO (18, 21).

By contrast, other researchers observe only a typical type 2 Cu(II) EPR signal for purified M. capsulatus (Bath) pMMO (14, 16), and for membrane-bound and whole cell suspensions of M. capsulatus (Bath) and Methylomicrobium album BG8 pMMO (22-24). Antholine and co-workers suggest that the isotropic signal attributed to a trinuclear cluster can instead be explained by the presence of a radical and a mononuclear Cu(II) ion coordinated by 3 or 4 nitrogen donors in a square planar geometry (22). In addition to copper, 1 to 2.5 iron ions have been detected in some preparations of purified pMMO from both M. capsulatus (Bath) (14, 16) and M. trichosporium OB3b (18). To address some of the many unresolved issues regarding pMMO, we have purified the enzyme from M. capsulatus (Bath), determined its molecular mass and metal ion content, and investigated the metal centers by X-ray absorption (XAS) and EPR spectroscopies.

Materials and Methods

Bacterial Growth.

M. capsulatus (Bath) was obtained from the American Type Culture Collection and from Professor Stephen J. Lippard (MIT). Cultures were maintained on agar plates containing potassium nitrate mineral salts (NMS) growth medium (25), 10 μ M CuSO₄ (26), and 0.25 mg/l Na₂MoO₄. Plates were stored in a BBL GasPak vented environmental chamber which was partially filled with methane gas (BOC Gases) and placed in a 42 °C incubator. Methane was replenished daily and plates restreaked every month. Inoculum for large scale fermentations was prepared by transferring a single colony to 1 ml NMS supplemented with 50 μ M CuSO₄ and 0.25 mg/l Na₂MoO₄ in a 25 ml culture tube. The tube was sealed with a septum, and 5 ml of the headspace evacuated and replaced with methane. After several days in a 42 °C shaking incubator at 250 rpm, the solution was subcultured into 5 ml and then into 50 ml of the above medium. The 50 ml flasks were recharged daily with methane and subcultured every third week. For 10 l fermentations, 50 ml cultures were added to 10 l NMS with 50 μ M CuSO₄ and 0.25 mg/l Na₂MoO₄ until an OD₆₀₀ of at least 0.2-0.4 was reached. Cells were grown at 42 °C by using a 1:4 methane:air mixture at 1 l/min and agitating at 300 rpm. When the OD_{600} reached 12-14, after approximately 48 hrs, cells were harvested by centrifugation at 4000 rpm for 20 min, washed three times with 10 mM Pipes buffer (pH 7.1-7.3), frozen in liquid nitrogen, and stored at -80 °C. Typically, 1 l of cells was retained in the fermenter as inoculum, and 9 l new growth medium was added.

Membrane Isolation and Protein Purification.

Cells were lysed by sonication, and cell debris removed by centrifugation at 12,000 rpm for 1 hr. The membrane fraction was isolated by ultracentrifugation for 1 hr at 40,000 rpm. Using a Dounce homogenizer, membranes were resuspended three times with 25 mM Pipes, 0.25 M NaCl (pH 7.2). After the final wash, the $\sim 10-20$ mg/ml membrane suspension was frozen in 1 ml aliquots in liquid nitrogen and stored at -80 °C. The detergent dodecyl-β-D-maltoside (12DM, Anatrace) was used to solubilize pMMO for purification (27). Membranes were thawed on ice and added to an equal volume of 50 mg/ml 12DM in ddH₂O. Before loading onto a Source 30Q (Amersham Biosciences) anion exchange column, solubilized membranes were diluted fourfold with the column equilibration buffer, 50 mM Hepes (pH 7.5), 0.05% 12DM. pMMO was eluted at 0.3-0.4 M NaCl using a 0-1 M NaCl gradient, concentrated in a Centriprep 50 (Millipore), and loaded onto a Sephacryl S200 gel filtration column (Amersham Biosciences) equilibrated with 50 mM Hepes, 0.25 M NaCl (pH 7.5), 0.05% 12DM. Purified pMMO was concentrated in a Centriprep 50 to 5-10 mg/ml and stored at 4 °C for immediate use or at -80 °C for future use. The concentration of solubilized pMMO was determined by the Detergent-Compatible (DC) Lowry assay (Bio Rad). Purified pMMO concentrations were determined using the extinction coefficient (585,016 M⁻¹cm⁻¹ per 200 kDa enzyme) determined by amino acid analysis (Protein Chemistry Laboratory, Texas A&M University).

Molecular Mass Determination.

SDS-PAGE (28), PFO-PAGE (29) and BN-PAGE (30) were performed as described previously with the following modifications. All samples were loaded onto gels immediately after adding sample buffers without boiling. Gradient gels (4-20%) for SDS-PAGE were obtained from ISC BioExpress. Tris/HCl gradient gels (4-15%, Bio Rad) were run on ice at 140 V for PFO-PAGE and on ice for 5 hr at 100 V for BN-PAGE. Coomassie Brilliant Blue G-250 for BN-PAGE was obtained from Sigma, and gels were destained for two days. Molecular mass markers typically used for analytical gel filtration (Amersham Biosciences) were used for PFO- and BN-PAGE as 1 mg/ml solutions in ddH₂O.

Activity Assays. Enzyme activity was measured by the epoxidation of propylene (31). In a typical experiment, 49 µl membrane-bound pMMO was mixed with 1 µl 1 mg/ml NADH in a 2 ml septum sealed serum vial (Wheaton). The reaction was initiated by evacuating 2 ml of air and adding 2 ml of propylene (BOC Gases). After shaking in a 45 °C water bath for 3 minutes, propylene oxide concentration was analyzed on a Hewlett Packard 5890 Gas Chromatograph with a Porapak Q column (Supelco). The amount of propylene oxide produced was determined by a standard curve using 1-10 mM propylene oxide (Aldrich) standards diluted in ddH₂O. The specific activity was calculated as the nmoles propylene oxide produced per minute per mg pMMO. To investigate the dependence of pMMO activity on metal ions, stock solutions of CuSO₄ and Na₂H₂EDTA were added to final concentrations of 2-200 µM Cu(II) and 0.2-72 uM EDTA. For solubilized and purified pMMO, duroquinol was substituted for NADH as a reductant (32). Duroquinone was reduced as described previously (14). All enzyme assays were performed in at least duplicate.

Metal Ion Content.

All glassware for metal ion analyses was acid washed for a minimum of 24 hours, rinsed thoroughly with chelexed ddH₂O, and dried. Metal ion content was measured by inductively coupled plasma-atomic emission spectroscopy (ICP-AES) using a Thermo Jarrell Ash Atomscan Model 25 Sequential ICP Spectrometer. Copper and iron atomic absorption standards (Aldrich) were prepared immediately before use at 2 ppm in chelexed ddH₂O for instrument calibration. In parallel with all protein samples,1 ppm copper and iron standards were measured. Prior to measurement, solubilized pMMO was diluted into chelexed ddH₂O.

CD and EPR Spectroscopies.

Circular dichroism (CD) spectra were measured in a 0.1 cm path length quartz cuvette using a Jasco J-715 spectrometer. To remove Hepes, which interferes with CD signals at low wavelengths, pMMO was exchanged twice into a 0.1% 12DM solution in ddH₂O using a Microcon 100 (Millipore). X-band EPR spectra were acquired at 120 K using a modified Varian E-4 spectrometer; Q-band EPR spectra were acquired at 2 K using an instrument described previously (33). EPR samples of membrane-bound and purified pMMO were frozen in liquid nitrogen. The concentration of EPR-active copper in pMMO was obtained by double integration using [CuEDTA]²⁻ as a standard. A 0.1 M [CuEDTA]²⁻ stock solution was prepared by adding equimolar CuCl₂ and Na₂H₂EDTA to a 1% NaOH solution. This solution was then serially diluted to 500 μM and 250 μM with ddH₂O.

X-ray Absorption Spectroscopy.

Purified samples for XAS were diluted in 50 mM Hepes (pH 7.5), 0.25 M NaCl, 0.05% 12DM, 50% glycerol and concentrated to 1-10 mM copper. Samples for reduction were vacuum degassed for 1.5 hr, transferred to a Coy anaerobic chamber, and chemically reduced by adding dithionite to a final concentration of 20 mM. Samples were loaded into Lucite cells wrapped with Kapton tape in air for as-isolated pMMO or anaerobically for reduced pMMO. Samples were flash frozen in liquid nitrogen and stored at -80 °C until data collection. XAS measurements were performed on reproducible independent samples.

Data were collected at beamline 7-3 at the Stanford Synchrotron Radiation Laboratory (SSRL) and at the Biophysics Collaborative Access Team (BioCAT) beamline at the Advanced Photon Source (APS) using Si(220) double crystal monochromators detuned 50% for harmonic rejection. Fluorescence excitation spectra at SSRL were recorded using a 30-element Ge solid-state array detector whereas APS data were collected using dual photo-multiplier detectors. Samples were maintained at 10 K using an Oxford Instruments continuous-flow liquid helium cryostat at SSRL and at 30 K using a He Displex Cryostat at APS. XAS spectra were measured at SSRL using 5 eV steps in the pre-edge, 0.25 eV steps in theedge (9636-9710 eV). and 0.05 Å⁻¹ increments in the extended X-ray absorption fine structure (EXAFS) region, integrating from 1s to 30s in a k^3 weighted manner for a total scan length of 40 minutes. X-ray absoprtion near edge structure (XANES) spectra were collected at APS using 0.25 eV point spacing at a constant 37 ms per point over an eV range of 8900 to 9100 for a total collection time of 30 seconds/scan. X-ray energies were calibrated by simultaneous measurement of a Cu foil absorption spectrum, assigning the first inflection point as 8980.3 eV. Each channel of every scan was examined for glitches prior to averaging, and spectra were closely monitored for photoreduction. SSRL data represent the average of 10 to 12 scans whereas APS data represent the average of 30 scans.

XAS data were analyzed using the Macintosh OS X version of the EXAFSPAK program suite integrated with Feff v7.0 for theoretical model generation (34, 35). Data reduction used a secondorder polynomial in the pre-edge and a three-region cubic spline through the EXAFS region. Data were converted to k-space using E₀=9000 eV. The k cubed weighted EXAFS was truncated at 1.0 and 13.0 Å⁻¹ for filtering purposes and Fourier transformed. Data were then Fourier-filtered to isolate the EXAFS for each peak in the Fourier transform. Fitting analysis was performed on both Fourier filtered and raw/unfiltered data. The best-fit bond lengths (R), coordination numbers (C.N.) and Debye-Waller factors (σ^2) listed in Table 3 were determined from fitting Fourier filtered (R=1.0-3.8 Å) data. Fits to both Fourier filtered and unfiltered data gave equivalent structural parameters,

however. EXAFS data were fit using amplitude and phase functions calculated with Feff v7.0 for a Cu-O at 2.00 Å, a Cu-C at 2.9 Å, a Cu-S at 2.25 Å and 2.40 Å, and a Cu-Cu at 2.60Å and 2.90Å. A scale factor of 0.86 and an E_0 of -12.2 were calibrated by fitting EXAFS data from crystallographically characterized complexes. Coordination numbers were fixed at half-integer values while R and σ^2 were freely variable parameters. The mean square deviation between data and fit, corrected for degrees of freedom, was used to determine the best simulation and justification for adding additional interactions to the simulation (36).

Results and Discussion

Isolation and Purification of pMMO.

Large quantities of *M. capsulatus* (Bath) cells were produced in 10 l fermentations using NMS medium supplemented with 50 μ M CuSO₄. In contrast to previous reports, high CuSO₄ concentrations did not inhibit cell growth (14, 17). It was therefore not necessary to gradually increase the copper levels (14) or to supplement the medium with CuEDTA (17, 20). Cells harvested at OD₆₀₀=12 yield ~10 g cell paste per liter medium. Membranes were washed thoroughly to remove adventitious metal ions and loosely bound proteins, and remain active after many months of storage at -80 °C. Reducing agents, catalase, and high ionic strength buffers (17) did not enhance stability.

Our purification protocol differs from those reported previously (14, 16, 17). The Source 30Q anion exchange column removes contaminating proteins as well as lipids. Low molecular mass proteins not resolved on the Source 30Q column are then separated by gel filtration on Sephacryl S200. The final pMMO sample is purified to homogeneity and comprises three polypeptides of molecular masses ~47 kDa, ~24 kDa, and ~22 kDa (Fig. 1*A*). A 35 kDa polypeptide, initially assigned as a pMMO subunit (26) and subsequently recognized to be a proteolytic fragment of the β subunit (17), is not observed even at high protein concentrations (Fig. 1*A*, lane 3). A 63 kDa putative formaldehyde or methanol dehydrogenase suggested to form a functional complex with pMMO (16) is typically removed prior to purification by washing the membranes. Samples of pMMO purified by this method are stable to proteolysis for many months at -80 °C or for 1-2 weeks at 4 °C.

Molecular Mass Determination. Although pMMO from *M. capsulatus* (Bath) is composed of three polypeptides, neither the subunit stoichiometry nor the total molecular mass has been determined. The ratio of subunits has always been assumed to be 1:1:1, corresponding to an $\alpha\beta\gamma$ heterotrimer of ~100 kDa. For *M. trichosporium* OB3b pMMO, comprising two subunits of molecular masses 25 and 41 kDa, the overall mass is estimated to be 326 kDa by analytical gel filtration (21). The molecular

mass of a multisubunit membrane protein cannot be measured accurately by analytical gel filtration chromatography, however (37). For this reason, other techniques, such as PFO-PAGE (29) and BN-PAGE (30), have been developed. These techniques are similar to SDS-PAGE except that the dodecylsulfate anion is replaced with a nondenaturing alternative that preserves the quarternary structure of the membrane protein.

PFO-PAGE analysis of purified pMMO reveals a band immediately below that of catalase (232 kDa) as well as one just above ovalbumin (43 kDa) (Fig. 1*B*). The ~200 kDa band suggests that purified pMMO has an $\alpha_2\beta_2\gamma_2$ subunit structure, although other compositions cannot be excluded. The lower molecular mass band likely corresponds to the α subunit (47 kDa), but could also be a complex between the β (24 kDa) and γ (22 kDa) subunits. PFO is thus partly denaturing under the experimental conditions. Analytical ultracentrifugation sedimentation velocity experiments using PFO running buffer also indicate the presence of two species corresponding to similar molecular masses (R. L. L., B. Demeler, A.C.R., unpublished results). A molecular mass of ~200 kDa is further corroborated by BN-PAGE (Fig. 1*O*). Lower molecular weight species are not observed in this experiment. Instead, a ~440 kDa molecular weight band is observed, perhaps corresponding to a tetramer. It is possible that pMMO can form higher order species considering the high levels of pMMO found in *M. capsulatus* (Bath) intracytoplasmic membranes (26). Most importantly, a ~100 kDa band is not apparent in either of these gel experiments. Since both techniques yield a ~200 kDa pMMO species, the amount of PFO or Brilliant Blue G-250 bound to the enzyme does not appreciably alter its mass.



Fig. 1. Gel electrophoresis analyses of purified pMMO. (A) SDS-PAGE. Lane 1, 1.5 mg/ml purified pMMO; lane 2, Invitrogen molecular mass standards: 116.3 kDa, 97.4 kDa, 66.3 kDa, 55.4 kDa, 36.5 kDa, 31 kDa, 21.5 kDa, 14.4 kDa, 6 kDa; lane 3, 6 mg/ml purified pMMO. (B) PFO-PAGE. Lane 1, ovalbumin (43 kDa); lane 2, albumin (67 kDa); lane 3, adenylate cyclase (70 kDa), lane 4, catalase (232 kDa); lane 5, urease monomer (91 kDa), lane 6: ferritin (440 kDa); lane 7, 2 mg/ml purified pMMO. (C) BN-PAGE. Lane 1, ovalbumin (43 kDa); lane 2, albumin (67, 134 kDa); lane 3, catalase (232 kDa); lane 4, ferritin (440 kDa); lane 5, 0.8 mg/ml purified pMMO.

Enzyme Activity.

The average specific activity of the membrane-bound pMMO is 19.4 nmol/mg min, comparable to other preparations (14, 16, 17, 19, 20) (Table 1). For membrane-bound pMMO, both NADH and duroquinol (32) were effective reducing agents. According to early reports, the addition of $CuSO_4$ either directly to membrane fractions (19) or to the cells prior to lysis and membrane isolation (38) resulted in increased pMMO activity. The activity of our membrane-bound pMMO was unchanged with the addition of up to 200 µM CuSO4, however. The cells for the previous experiments were grown in the presence of only 0-20 μ M CuSO₄, which could lead to incomplete metallation of the pMMO active site. In the present experiments, the cells were grown with 50 μ M CuSO₄ and may already contain fully loaded active sites. The effect of EDTA on pMMO activity was also investigated. The addition of 0.2-72 mM EDTA decreased activity, but approximately half of the original specific activity remained even with a 40-fold molar excess of EDTA. A similar effect has been observed for purified pMMO from *M. trichosporium* OB3b (18) and for partially purified pMMO from *M. capsulatus* (Bath) (16). These observations suggest that the metal ions in pMMO are necessary for activity, but may differ in accessibility. The specific activity decreased upon solubilization with 12DM (Table 1). No activity was observed when NADH was used as the reductant, but was detected with duroquinol. Attempts to correlate solubilized pMMO activity with detergent levels did not yield an optimal detergent-to-protein ratio. After purification, most samples were not active. The 17.7 nmol/mg min activity measured for the few active preparations (Table 1) is higher than the values reported for other purified samples (14, 17). The ability to detect activity could not be correlated with metal ion content or with particular protein impurities.

Table 1. Average specific activity of pMMO								
pMMO sample	Reductant	Specific activity (nmole/min mg)						
Membrane- bound	NADH	21						
	duroquinol	16						
Solubilized	duroquinol	3.9						
Purified	duroquinol	17.7						

For inactive samples, numerous approaches to restore activity were pursued. Attempts to remove bound detergent by dialysis, to add back other proteins separated on the anion exchange column, and to supplement the assay mixture with reducing agents or CuSO₄ and CuCl were not successful. A lipid reconstitution procedure reported to yield recovered activities of 2-5 nmol/mg min (17) did not restore activity either. There are several possible explanations for the lack of activity. First, the detergent could interfere with activity. Second, a cofactor or protein essential to the electron transport chain might be removed during purification. Potential candidates include cytochromes associated with pMMO (14, 17), an NADH: quinone oxidoreductase (39), or a putative dehydrogenase suggested to form an active complex with pMMO (16). Finally, solubilized, purified pMMO might not be folded properly. The CD spectrum shows secondary structure with a significant α -helical component (Fig. 2), but it is still possible that pMMO is present in an inactive conformation. Until these issues are fully investigated, the variability in pMMO activity levels observed in our and other preparations (17) will likely remain a problem.



Fig. 2. CD spectrum of 0.77 μ M purified pMMO in 12DM. Data were collected at room temperature from 300 to 190 nm at 1 nm resolution

Metal Ion Analyses. According to ICP-AES measurements, membrane-bound pMMO contains 20.8 \pm 4.5 copper ions per 200 kDa, and purified pMMO contains 4.8 ± 0.8 copper ions per 200 kDa (Table 2). By contrast, values of 8-118 copper ions per 200 kDa for membrane-bound pMMO (14, 19, 20, 38) and 24-30 copper ions per 200 kDa for purified pMMO (14, 17) have been reported. There are several likely explanations for the presence of significantly fewer copper ions in our pMMO samples.

First, the wide variation in membrane-bound pMMO copper content reported by others suggests that cells and membranes may not have been washed thoroughly. Large amounts of copper present in the growth medium could remain adventitiously bound to the membranes, and could also account for the high levels of copper found in purified pMMO. For our preparations, both the harvested cells and membrane fractions were washed carefully three times. Second, the protein concentration may not have been determined accurately, leading to incorrect stoichiometry. For example, we found that the Bradford assay is inaccurate in the presence of 12DM. Thus, for our ICP-AES analysis, the concentration of solubilized pMMO was determined using a detergent-compatible version of the Lowry assay, and the concentration of purified pMMO was calculated using an extinction coefficient determined from amino acid hydrolysis. In support of our results, Dalton and co-workers also measured 4 copper ions per 200 kDa using a detergent-compatible protein assay (16). Although it is possible that our purification procedure removes some essential copper ions, the activity is in fact similar to that observed for preparations containing significantly more copper (14, 19, 20). Thus, it seems likely that the catalytic metal center(s) remain(s) intact.

Although copper is always detected in pMMO, the presence of iron has been disputed (17). For membrane-bound pMMO, iron stoichiometries ranging from 2 to 20 iron ions per 200 kDa have been reported (14, 19, 20). For purified pMMO, values of 0 (17), 2 (16), or 5 (14) iron ions per 200 kDa have been reported. Our membrane-bound pMMO contains 2.2 ± 0.7 iron ions per 200 kDa, and our purified pMMO contains 1.5 ± 0.7 iron ions per 200 kDa (Table 2). Chan and co-workers do not detect iron in their purified pMMO and suggest that high iron concentrations in the growth medium lead to pMMO contamination by sMMO and other iron-containing proteins (17, 20). Although our cell growth medium includes 80 μ M FeEDTA, sMMO is not associated with our membrane-bound or purified samples. The small amount of iron in our purified samples may instead be due to cytochrome impurities. Nevertheless, we cannot exclude the possibility that iron plays a functional role in pMMO.

Table 2. Metal ion analysis of pMMO							
pMMO sample	Mol Cu/ 200 kDa	Mol Fe/200 kDa					
Solubilized*	20.8 ± 4.5	2.2 ± 0.7					
Purified [#]	4.8 ± 0.8	1.5 ± 0.7					

^{*}Average of 7 samples †Average of 28 samples

EPR Spectroscopy.

The Q-band (35 GHz) EPR spectra of both membrane-bound and purified pMMO exhibit signals characteristic of type 2 Cu(II) with $g_{11} = 2.24$, $g_{\perp} = 2.05$, and $A_{11} = 530$ MHz (Fig. 3). These parameters are similar to those reported for both whole cell suspensions (24) and purified samples (14, 16) from *M. capsulatus* (Bath), and are consistent with a mononuclear Cu(II) site coordinated by 3 or 4 histidine imidazoles (22-24, 40). By contrast, the signal attributed by Chan and co-workers to a mixed valence trinuclear copper cluster (17, 19, 20) is not observed in either spectrum. A radical at g = 2.00 and a signal from trace amounts of rhombic iron are observed in the membrane-bound sample, but are not observed upon purification. Integration of the X-band EPR spectrum indicates that 40-60% of the total copper (4-6 copper ions per 200 kDa) in purified pMMO are EPR-active. An increase in signal intensity is not obtained by adding ferricyanide, in contrast to previous reports (20). Upon addition of 20 mM dithionite, the Cu(II) EPR signal disappears completely.



Fig. 3. Q-band EPR spectra of purified (top) and membrane-bound (bottom) pMMO. The same signal is observed for both samples with g₁₁ = 2.24, g₁ = 2.05, and A₁₁ (Cu) = 530 MHz. Spectrometer conditions: microwave frequency, 35.72 GHz; microwave power, 50 μW; modulation amplitude, 1.3 G; scan time, 4 min; time constant, 128 msec; temperature, 2 K.



Fig. 4. Cu K-edge X-ray absorption near edge spectra of purified pMMO. Dashed line, as isolated purified pMMO; solid line, dithionite reduced pMMO. The 1s \rightarrow 4p feature at ~8984 eV is marked by the dashed vertical line.

X-Ray Absorption Spectroscopy.

XANES spectra for purified and purified dithionite reduced pMMO are compared in Fig. 4. A substantial (~3 eV) edge energy shift observed upon chemical reduction indicates that the average copper oxidation state of as-isolated pMMO is higher than that of the dithionite reduced sample. An absorption feature at ~8984 eV, attributed to a Cu(I) 1s \rightarrow 4p transition (41), is present in both spectra, however. Therefore, some fraction of the copper in both pMMO samples is in the Cu(I) oxidation state, and this fraction is clearly greater for the reduced protein. The presence of Cu(I) in the as-isolated purified protein is consistent with the EPR quantitation. Previous XANES spectra of pMMO membrane preparations also suggested a mixture of copper oxidation states, although the ~8984 eV feature was less pronounced upon reduction (20).

Table 3. Summary of pMMO EXAFS fitting analysis*												
fit no.		1			2			3			4	
	C.	R (Å)	σ^2 (Ų)†	C.N	R (Å)	σ^2 (Å ²)†	C.N	R (Å)	σ^2 (Å ²)†	C.N	R (Å)	σ² (Ų)†
	N.									•		
Cu-	2.5	1.97	3.17	2.5	1.97	3.02	2.5	1.97	2.82	2.5	1.97	3.70
N/O												
Cu-C				1.0	2.60	1.70						
Cu-S							1	2.71	3.13			
Cu-Cu										0.5	2.57	3.73
F'‡		271			69			59.6			40.7	

*The best fit is listed in bold.

[†]Debye-Waller factor x 10³.

[‡]Degrees for freedom weighted mean square deviation between data and fit.

The Fourier transform of the Cu EXAFS data (Fig. 5) exhibits two peaks, indicating that two shells of backscatterers are present. The EXAFS generating the more intense peak at 1.5 Å are best fit with oxygen/nitrogen (O/N) ligands at average bond lengths of 1.97 Å (Table 3, fit 1). The corresponding coordination number is 2.5 O/N ligands per copper ion, but this value may be underestimated. Cu-O/N bond lengths of ~2 Å are typically observed for four-coordinate Cu(I) and Cu(II) species, whereas two- and three-coordinate Cu-O/N bond lengths are on average 0.15-0.06 Å shorter, respectively (42). Two scenarios would result in an underestimate of the Cu-O/N coordination number. First, the low coordination number may reflect the presence of two sets of Cu-O/N ligand environments at close but distinct bond lengths. Two such environments cannot be fit independently given the data resolution. Alternatively, a low percentage of Cu-S scattering might destructively interfere with the Cu-low Z signal and dampen the EXAFS amplitude for the Cu-O/N interaction. There is no direct XAS evidence for a Cu-S interaction, however.

The second peak in the Fourier transform, observed reproducibly at a phase shifted bond length of 2.2 Å, corresponds to a second backscattering shell at ~2.6 Å (Fig. 5.4). Attempts to fit this feature employed three models: a Cu-C interaction, possibly from a nearby histidine imidazole ring (Table 3, fit 2), a long Cu-S interaction (Table 3, fit 3), and a direct Cu-Cu interaction (Table 3, fit 4). In each case, addition of a second ligand shell produced a mean-squared deviation value (F') better than a single environment fit (Table 3, fit 1), but the interaction was best simulated by a second Cu atom at 2.57 Å (Fig. 2*B*). Notably, the 2.57 Å Cu-Cu distance is strikingly similar to Cu-Cu distances observed in the nitrous oxide reductase tetranuclear Cu_Z cluster (43). Short Cu-Cu distances are also present in the cytochrome *c* oxidase Cu_A site (44), but optical and EPR spectroscopic features characteristic of Cu_A (45) or Cu_Z (46) are not observed for pMMO. Thus, pMMO may contain a new type of copper center. Although it is likely that the second metal ion in pMMO is copper, further analysis is required to rule out the possibility that another metal ion, such as iron, is involved. Nevertheless, this result provides the first direct evidence that some component of the pMMO metal ions exist in a multinuclear, copper-containing cluster.



Fig. 5. Cu EXAFS fitting analysis for pMMO. (A) Fourier transform comparison of raw/unfiltered pMMO EXAFS (solid line) with two environment Cu-O/N and Cu-Cu theoretical fit (dashed line). (B) Comparison of unfiltered k³ weighted pMMO EXAFS (solid line) with a theoretical two environment Cu-O/N and Cu-Cu k³ weighted EXAFS (dashed line).

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