

## Title

**Enzymatic Oxidation of Methane**

## authors

Sarah Sirajuddin and Amy C. Rosenzweig\*

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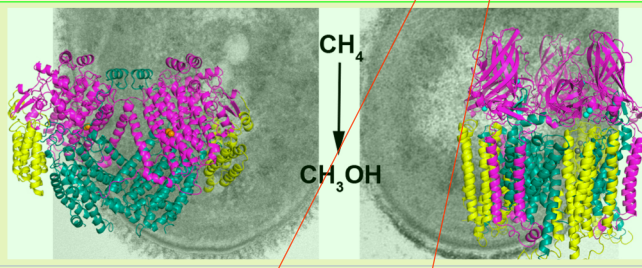
Departments of Molecular Biosciences and of Chemistry, Northwestern University, Evanston, Illinois 60208, United States

## Content

**ABSTRACT:** Methane monooxygenases (MMOs) are enzymes that catalyze the oxidation of methane to methanol in methanotrophic bacteria. As potential targets for new gas-to-liquid methane bioconversion processes, MMOs have attracted intense attention in recent years. There are two distinct types of MMO, a soluble, cytoplasmic MMO (sMMO) and a membrane-bound, particulate MMO (pMMO). Both oxidize methane at metal centers within a complex, multisubunit scaffold, but the structures, active sites, and chemical mechanisms are completely different. This Current Topic

review article focuses on the overall architectures, active site structures, substrate reactivities, protein–protein interactions, and chemical mechanisms of both MMOs, with an emphasis on fundamental aspects. In addition, recent advances, including new details of interactions between the sMMO components, characterization of sMMO intermediates, and progress toward understanding the pMMO metal centers are highlighted. The work summarized here provides a guide for those interested in exploiting MMOs for biotechnological applications.

## Image



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Aerobic methane oxidation in nature is carried out by methanotrophic bacteria, organisms that can use methane as their sole source of carbon and energy. Despite the extremely strong C–H bond in methane ( $\Delta H_{298} = 105 \text{ kcal/mol}$ ),<sup>1</sup> these bacteria are capable of oxidizing methane to methanol under ambient conditions using metalloenzymes called methane monooxygenases (MMOs). Understanding the biochemistry of MMOs is critical to their applications in bioremediation<sup>2</sup> and to the development of biological processes for conversion of natural gas to liquid fuels and chemicals.<sup>3</sup> Moreover, detailed characterization of MMO active sites may lead to new biomimetic catalysts.<sup>4</sup> A range of alkane-oxidizing enzymes exist in the environment,<sup>5</sup> but only the two types of MMO, soluble MMO (sMMO) and particulate MMO (pMMO), as well as the close pMMO homologue ammonia monooxygenase (AMO),<sup>6</sup> can oxidize methane. Although sMMO and pMMO perform the same chemical reaction, their structures, cofactor requirements, and mechanisms are completely different.

sMMO requires three protein components for activity: a hydroxylase (MMOH, 251 kDa), a reductase (MMOR, 38.5 kDa), and a regulatory protein (MMOB, 15.9 kDa).<sup>7</sup> MMOH is responsible for methane oxidation and is composed of  $\alpha$ ,  $\beta$ , and  $\gamma$  subunits encoded by the *mmoX*, *mmoY*, and *mmoZ* genes, respectively. MMOB and MMOR are encoded by the *mmoB* and *mmoC* genes. The pMMO hydroxylase consists of the *pmoB*, *pmoA*, and *pmoC* subunits, encoded by the *pmoCAB* operon (Figure 1).<sup>8</sup> No regulatory protein has been identified for pMMO, and the identity of its native reductase remains unclear.<sup>9</sup> Most methanotrophs express only pMMO. A few species express only sMMO, and a subset contains genes for both pMMO and sMMO.<sup>10</sup> In this subset, sMMO expression occurs when copper levels in the growth medium are below 1  $\mu\text{M}$ , and pMMO expression is predominant at copper levels

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above 5  $\mu\text{M}$ .<sup>11–13</sup> This “copper switch” parallels the metal cofactor requirements of the MMOs: sMMO has a catalytic diiron center,<sup>14</sup> and pMMO utilizes a copper active site.<sup>9</sup> The mechanism of the copper switch is still unknown, but recent work suggests that the copper-chelating natural product methanobactin, genes encoded by the methanobactin operon, and *mmoD*, a gene product of unknown function encoded by the sMMO operon, could play a role.<sup>15,16</sup> A transcriptional activator, *mmoR*, and a GroEL-like chaperone protein, *mmoG* (Figure 1), are required for sMMO expression.<sup>17,18</sup>

Studies of both MMOs over the past quarter century have provided much insight into their structure and function. This Current Topic summarizes work addressing the architecture, active sites, substrate reactivities, protein–protein interactions, and chemical mechanisms of both systems, with emphasis on fundamental aspects that must be considered if MMOs are to be exploited in new gas-to-liquid methane bioconversion processes. In addition, recent advances that have not been reviewed previously are highlighted. A more comprehensive review is available elsewhere.<sup>19</sup>

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**ENZYME ARCHITECTURE**

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**Soluble MMO.** To date there are 28 crystal structures of MMOH, primarily from *Methylococcus capsulatus* (Bath), one structure of the MMOH–MMOB complex, two NMR structures of MMOB, and two NMR structures of MMOR (Table 1). The first crystal structure of MMOH was determined in 1993, and a resolution of 1.7 Å has been achieved.<sup>20,21</sup> The  $\alpha$  (60.6 kDa),  $\beta$  (45.1 kDa), and  $\gamma$  (19.8

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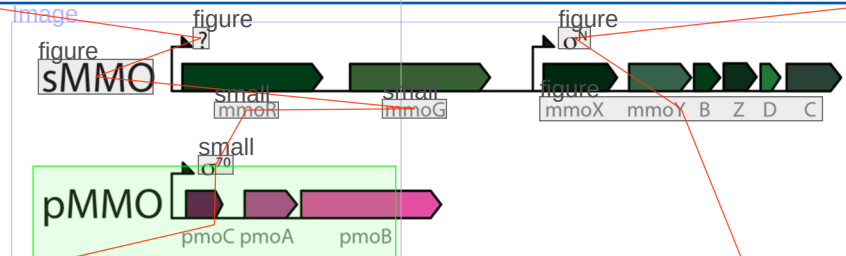
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Figure 1. sMMO and pMMO operons.

table

**Table 1. MMO Structures**

	small resolution [Å]	small PDB code
<i>Methylococcus capsulatus</i> (Bath) MMOH (hydroxylase)		
oxidized 4 °C	2.20	1MTY
small	2.20	1MTY
small	2.16	1FZ1
small	2.10	1MTY
small	2.15	1FYZ
small	2.40	1FZS
small	2.15	1FZ2
small	2.07	1FZ0
methanol soaked	2.05	1FZ6
small		
ethanol soaked	1.96	1FZ7
datum	3.30	1FZ1
Xe pressurized		
datum	2.60	1EZB
dibromomethane grown	2.10	1EZ8
iodoethane grown	2.30	1EZ9
pH 8.5 soaked	2.38	1EZ4
pH 6.2 soaked	2.03	1FZ3
small	2.32	1XMF
Mn(II) soaked		
apo (metal free)	2.10	1XMG
Cu(II) reconstituted	2.32	1XMH
phenol soaked	1.96	1XUS
6-bromohexanol soaked	1.80	1XVB
8-bromoocanol soaked	2.00	1XVC
4-fluorophenol soaked	2.30	1XVD
3-bromo-3-butanol soaked	2.40	1XVE
chloropropanol soaked	2.00	1XVR
datum	1.96	1XVG
bromothanol soaked	small	1XU1
bromophenol soaked	2.30	1XU3
<i>Methylosinus trichosporium</i> OB3b MMOH (hydroxylase)		
oxidized	small	1MTH
small	2.00	1MTH
oxidized	2.70	1MTH
MMOB, MMOC, and Protein-Protein Complexes		
<i>M. capsulatus</i> (Bath) MMOB NMR		1CKV
<i>M. trichosporium</i> OB3b MMOB NMR		2MOB
<i>M. capsulatus</i> (Bath) MMOC [2Fe-2S] domain		1JQ4
NMR	small	1TVQ
<i>M. capsulatus</i> (Bath) MMOC FAD/NADH binding domain NMR		1YEW
<i>M. capsulatus</i> (Bath) MMOH-MMOB complex	small	4GAM
<i>M. capsulatus</i> (Bath) pMMO	small	1YEW
<i>M. capsulatus</i> (Bath) pMMO	small	3RGB <sup>a</sup>
<i>M. trichosporium</i> OB3b pMMO	small	3CHX
<i>Methylocystis</i> species strain M pMMO	small	3RER
<i>Methylocystis</i> species strain Rockwell pMMO	small	3PHZ
<i>Methylocystis</i> species strain Rockwell pMMO Cu(II) soaked	3.15	4PI0
<i>Methylocystis</i> species strain Rockwell pMMO Zn(II) soaked	3.33	4PI2

other

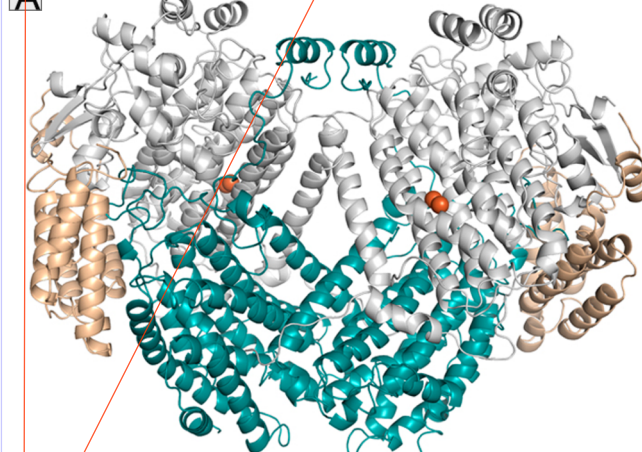
"PDB 3RGB is an improved version of structure 1YEW and should be used as the *M. capsulatus* (Bath) pMMO model; 1YEW is obsolete."

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kDa) subunits are arranged as an  $\alpha_2\beta_2\gamma_2$  dimer composed mainly of  $\alpha$ -helices except for two  $\beta$  hairpins in the  $\alpha$  and  $\gamma$  subunits (Figure 2A). The dimer interface involves interactions between the  $\alpha$  and  $\beta$  subunits, and a canyon region formed by

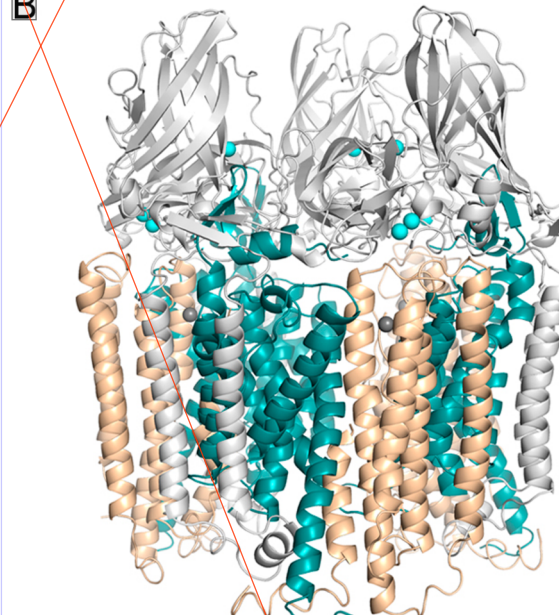
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A



figure

B



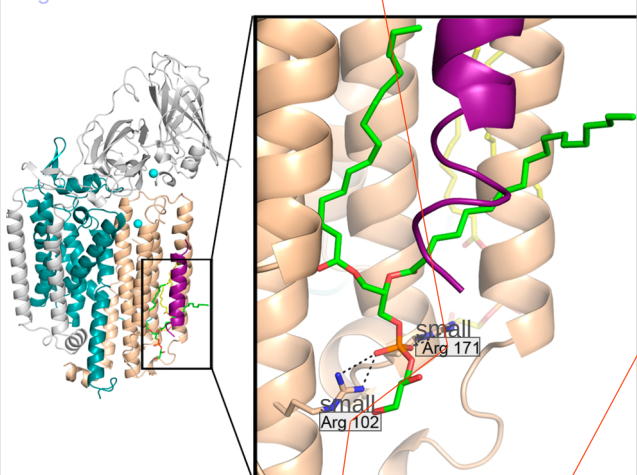
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Figure 2. Overall architecture of MMOs. (A) The sMMO hydroxylase (MMOH, PDB accession code 1MTY) with  $\alpha$  subunits shown in gray,  $\beta$  subunits shown in teal, and  $\gamma$  subunits shown in wheat. Each  $\alpha_2\beta_2\gamma_2$  dimer contains two diiron active sites (orange spheres). (B) The pMMO trimer (PDB accession code 3RGB) with pmoB subunits shown in gray, pmoA subunits shown in teal, and pmoC subunits shown in wheat. Copper ions are shown as cyan spheres and zinc ions are shown as gray spheres.

the  $\alpha$  and  $\beta$  subunits in each  $\alpha\beta\gamma$  protomer provides the docking surface for MMOR and MMOB. MMOB, which has no cofactors, consists of seven  $\beta$  strands arranged in two  $\beta$  sheets and three  $\alpha$  helices. MMOR consists of two mixed  $\alpha\beta$  domains: an FAD- and NADH-binding domain provides electrons to a [2Fe-2S] ferredoxin domain for transfer to the MMOH active site.<sup>7,22</sup>

**Particulate MMO.** The first crystal structure of pMMO was that of the *M. capsulatus* (Bath) enzyme, determined in 2005 to 2.8 Å resolution.<sup>23</sup> Structures of pMMOs from *Methylosinus trichosporium* OB3b, *Methylocystis* species strain M, and *Methylocystis* species strain Rockwell have also been reported (Table 1).<sup>24–26</sup> The pmoB, pmoA, and pmoC subunits are arranged as a 300 kDa  $\alpha_3\beta_3\gamma_3$  trimer (Figure 2B), which is consistent with electron microscopic data.<sup>27</sup> The pmoB subunit (42 kDa) consists of two cupredoxin-like domains linked by two transmembrane helices. The pmoA (24 kDa) and pmoC (22 kDa) subunits are composed primarily of transmembrane helices. In three of the four pMMO crystal structures, an additional helix not connected to any of the pMMO subunits is observed and appears to interact closely with the pmoC subunit.<sup>24–26</sup> In the *Methylocystis* sp. str. Rockwell pMMO structure, this interaction is mediated by a lipid molecule (Figure 3).<sup>26</sup> Since pMMO is crystallized from the native organism, this yet to be identified helix is likely physiologically relevant.

Image



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Figure 3. One  $\alpha\beta\gamma$  protomer of *Methylocystis* sp. strain Rockwell pMMO (PDB accession code 4PHZ) showing the unidentified helix (purple) and two bound lipids (green and yellow). The unidentified helix is also observed in structures of pMMO from *M. trichosporium* OB3b and *Methylocystis* sp. strain M. The lipid shown in green forms hydrogen bonds with conserved arginine residues from pmoC and mediates the interaction between pmoC and the mystery helix.

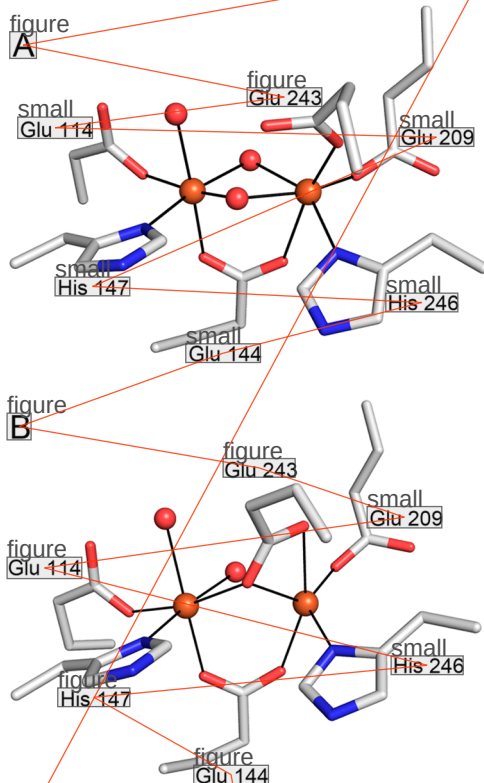
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## METAL CENTERS AND ACTIVE SITE STRUCTURE

**Soluble MMO.** MMOH contains only one type of metal center, the diiron active site located in each  $\alpha$  subunit in the  $\alpha_2\beta_2\gamma_2$  dimer. This diiron center is housed within a four-helix bundle of which surface-exposed helices E and F play key roles in modulating structural changes throughout the catalytic cycle.<sup>22</sup> Fe1 is coordinated by His147, Glu114, and a water molecule, and Fe2 is coordinated by His246, Glu243, and Glu209.<sup>20</sup> The two iron ions are bridged by residue Glu144 and two hydroxides (Figure 4A). Upon reduction, the Fe–Fe

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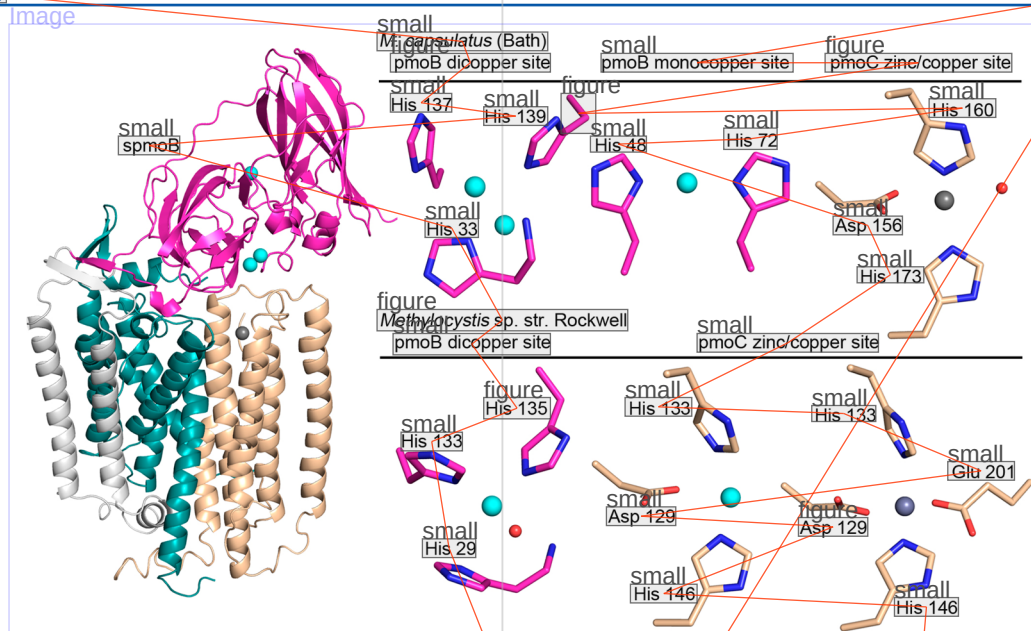
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Figure 4. Active site of sMMO. (A) Oxidized diiron center (PDB accession code 1MTY). (B) Reduced diiron center (PDB accession code 1FYZ). Iron ions are shown as orange spheres and solvent ligands are shown as red spheres.

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distance increases from ~3.1 Å to ~3.3 Å, and Glu243 undergoes a carboxylate shift, replacing the bridging hydroxide ion as a ligand to the diiron center. This shift opens a site for reaction with dioxygen (Figure 4B).<sup>21,28</sup> It is well established that the two iron ions are present as an antiferromagnetically coupled diiron(III) site in oxidized MMOH and a weakly ferromagnetically coupled diiron(II) center in reduced MMOH.<sup>29–32</sup> A mixed valence form can also be generated, but does not play a role in the catalytic cycle.<sup>7</sup>

**Particulate MMO.** In contrast to MMOH, several metal centers are observed in the structures of pMMO. All structures contain a copper binding site at the N-terminus of the pmoB subunit with ligands His33, which is the amino terminal residue of pmoB, His137, and His139 (*M. capsulatus* (Bath) numbering) (Figure 5). Guided by extended X-ray absorption fine structure (EXAFS) data,<sup>33</sup> this site was modeled as two copper ions with a Cu–Cu distance of ~2.5 Å in the structures of *M. capsulatus* (Bath)<sup>23</sup> and *M. trichosporium* OB3b<sup>24</sup> pMMO, and in one protomer of *Methylocystis* sp. str. M pMMO.<sup>25</sup> However, the moderate crystallographic resolution (2.6–3.9 Å) precludes definitive assignment of the nuclearity. This site was modeled as monocupper in *Methylocystis* sp. str. Rockwell pMMO<sup>26</sup> and in two protomers of *Methylocystis* sp. str. M pMMO.<sup>25</sup> Both of these pMMOs exhibit low copper binding stoichiometries once the enzyme is purified and concentrated. In the *Methylocystis* sp. str. Rockwell pMMO structure, a solvent molecule is modeled coordinated to the single copper ion (Figure 5) and is within hydrogen bonding distance of Glu31 (Glu35 using *M. capsulatus* (Bath) numbering).<sup>26</sup> A monocupper site is also present at the equivalent location in the 1.8 Å resolution structure of the soluble domain of the amoB



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**Figure 5.** Metal centers in the pMMO crystal structures. The locations of the dicopper, monocopper, and zinc/copper sites are shown as cyan and gray spheres within a protomer of *M. capsulatus* (Bath) pMMO (left). In the spmoB protein, two transmembrane helices (gray) linking the periplasmic N- and C-terminal cupredoxin domains of pmoB (magenta) are replaced with a GKLGGG linker connecting residue 172 to residue 265. The structure of spmoB has not been determined. The *M. capsulatus* (Bath) pMMO structure contains dicopper, monocopper, and zinc sites (top right, PDB accession code 3RGB). In the *Methylocystis* sp. str. Rockwell pMMO structure (bottom right), the dicopper site is modeled with a single copper ion and a solvent molecule, and the zinc/copper site is occupied by copper (PDB accession code 4PHZ). Soaking the crystals in zinc reveals a fourth ligand to this site, Glu201 (PDB accession code 4PI2). Solvent ligands are shown as red spheres.

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subunit from *Nitrosocaldus yellowstonii* AMO, indicating conservation of this site over the broad pMMO/AMO superfamily. However, in this structure, the copper ion is involved in a crystal lattice contact, and the N-terminal histidine ligand is disordered.<sup>34</sup> In the *M. capsulatus* (Bath) pmoB subunit, an additional copper ion is present at the interface of the two cupredoxin domains, coordinated by His48 and His72 (Figure 5). The His48 ligand is not conserved in other pMMO sequences,<sup>35</sup> and the site is not observed in the other pMMO structures.<sup>24–26</sup>

Several lines of evidence suggest that the crystallographically modeled dicopper center is the active site of pMMO. First, the ligands to the dicopper center are conserved in almost all methanotroph pmoB sequences as well as in homologous subunits from all members of the pMMO/AMO superfamily.<sup>34</sup> The only exceptions are a few pmoB sequences from methane-oxidizing Verrucomicrobia.<sup>36,37</sup> These sequences are evolutionarily distant, and it is possible that their extreme growth environments (pH 0.8–5, 22–88 °C) require a different type of active site.<sup>9</sup> Second, metal loading of apo membrane-bound pMMO only results in methane oxidation after addition of 2–3 copper equivalents per 100 kDa  $\alpha\beta\gamma$  pMMO protomer, consistent with an active site requiring more than one copper ion.<sup>26,38</sup> Third and most important, a recombinant soluble fragment of the *M. capsulatus* (Bath) pmoB subunit (spmoB, Figure 5) can oxidize methane to methanol in the presence of copper, and this activity is abolished upon mutagenesis of two or all three of the ligands to the dicopper site.<sup>38,39</sup> Consistent with this model, mutagenesis of the ligand equivalent to His139 in a homologous *Mycobacterium* NBB4 hydrocarbon monooxygenase (HMO) that oxidizes C2–C4 alkanes reduces activity significantly. However, in that enzyme, ~20% residual activity is retained,<sup>40</sup> perhaps due to copper binding by the other ligands, as has been observed for spmoB.<sup>39</sup>

Both Cu<sup>I</sup> and Cu<sup>II</sup> are routinely detected in pMMO by X-ray absorption spectroscopy,<sup>33</sup> and all samples of pMMO exhibit an electron paramagnetic resonance (EPR) spectroscopic signal attributable to a type 2 Cu<sup>II</sup> site.<sup>39,41,42</sup> Correlation of EPR signals with specific crystallographic copper sites in *M. capsulatus* (Bath) pMMO has been facilitated by comparative analysis of pMMO and the spmoB proteins.<sup>39</sup> EPR and metal analysis using pMMO and spmoB variants that disrupt the monocopper (H48N,H72A), dicopper (H33,137,139A), or both copper (H48N\_H33,72,137,139A) sites indicate the presence of a valence localized, mixed valence Cu<sup>I</sup>Cu<sup>II</sup> center at the dicopper site and a Cu<sup>I</sup> ion at the monocopper site. The location of the Cu<sup>II</sup> is proposed to be valence scrambled between the two copper binding sites with one copper coordinated by His137 and His139, and one coordinated by His33 and the N-terminal amino group. No water or hydroxide ligands to the Cu<sup>II</sup> ion are detected.

An additional metal binding site is present in the pmoC subunit of pMMO. This site can bind zinc or copper, and the metal ion is coordinated by residues Asp156, His160, and His173 (*M. capsulatus* (Bath) numbering) (Figure 5).<sup>23–26</sup> Recent metal binding studies using pMMOs from *M. capsulatus* (Bath) and *Methylocystis* sp. str. Rockwell suggest that zinc binding at this site partially inhibits activity, perhaps by interfering with proton transfer, and that this site is likely unoccupied by metal ions in vivo.<sup>26</sup> Interestingly, a structure of *Methylocystis* sp. str. Rockwell pMMO soaked in zinc reveals a fourth ligand to this site, Glu201 (Glu 228 using *M. capsulatus* (Bath) numbering) (Figure 5), which derives from a loop region of pmoC that is disordered in all previous structures. This region spanning pmoC residues 200–223 (*Methylocystis* sp. str. Rockwell numbering) contains many strictly invariant residues, including Glu211, Phe213, His218, Phe221, and Val222 (Glu238, Phe240, His245, Phe248, and Val249 using *M.*

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*capsulatus* (Bath) numbering). The potential importance of this pmoC region is underscored by mutagenesis studies of the *Mycobacterium* NBB4 HMO in which individual mutations in the residues corresponding to ligands Asp156, His160, and His173 (*M. capsulatus* (Bath) numbering) completely abrogated hydrocarbon oxidation activity.<sup>40</sup> Replacement of the residue equivalent to Asp166 (*M. capsulatus* (Bath) numbering) also abolished activity, which is interesting because this residue is located on a flexible loop that points away from the metal binding site in the pMMO structures.

**heading****SUBSTRATE BINDING AND REACTIVITY****Content**

**Soluble MMO.** In most studies, the activity of sMMO has been determined by measuring the epoxidation of propylene. This assay was preferred historically because the product, propylene oxide, is not further oxidized by methanotroph cells and extracts.<sup>43</sup> For *M. capsulatus* (Bath) sMMO, the turnover number for methane is in the range of  $0.2\text{--}1.0\text{ s}^{-1}$  at  $45^\circ\text{C}$  and that for propylene is  $\sim 0.7\text{ s}^{-1}$  (Table 2).<sup>44,45</sup> The  $K_m$  values for

**figure****Table 2. MMO Turnover Numbers**

		sample	organism	turnover number per active site ( $\text{s}^{-1}$ )	reference
cells producing sMMO	<i>M. trichosporium</i> OB3b	>3.5		10, 46	137
cells producing pMMO	<i>M. trichosporium</i> OB3b	0.5–2.5 <sup>a</sup>		66	66
	<i>M. capsulatus</i> (Bath)	small	0.5–2.5 <sup>a</sup>	12, 67	12, 67
purified sMMO	methane	<i>M. capsulatus</i> (Bath)	0.2–1.0	44, 45	
		<i>M. trichosporium</i> OB3b	3.7	46	
	propylene	<i>M. capsulatus</i> (Bath)	0.7	44, 45	
		<i>M. trichosporium</i> OB3b	4.4	46	
membrane-bound pMMO	methane	<i>M. capsulatus</i> (Bath)	0.05–0.27 <sup>b</sup>	26	
	propylene	<i>M. capsulatus</i> (Bath)	0.033–0.25	12, 65	
		<i>M. trichosporium</i> OB3b, <i>Methylocystis</i> sp. str. M, <i>Methylocystis</i> sp. str. Rockwell	0.003–0.04	9, 26	
purified pMMO	methane		NA		
	propylene	<i>M. capsulatus</i> (Bath)	0.003–0.27	10	

<sup>a</sup>Assuming cells are 50% protein and MMO is 20% of total protein.

<sup>b</sup>Assuming membrane samples are 80% pMMO.

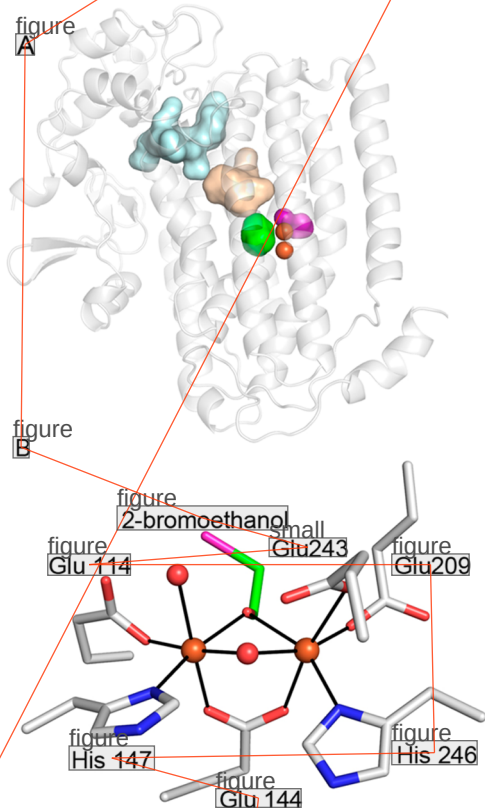
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methane and propylene are 0.94 and 2  $\mu\text{M}$ , respectively, and the  $K_m$  value for  $\text{O}_2$  is  $\sim 16\text{ }\mu\text{M}$  with both substrates.<sup>44</sup> For *M. trichosporium* OB3b sMMO, turnover numbers of  $3.7\text{ s}^{-1}$  and  $4.4\text{ s}^{-1}$  at  $30^\circ\text{C}$  have been reported for methane and propylene, respectively,<sup>46</sup> and the  $K_m$  for methane is  $\sim 12\text{ }\mu\text{M}$ .<sup>47</sup> The whole cell *M. trichosporium* OB3b sMMO turnover number is at least  $3.5\text{ s}^{-1}$  (Table 2).<sup>46</sup>

sMMO can oxidize a wide range of hydrocarbons, including C1–C8 *n*-alkanes, alkenes, and larger substrates such as benzene, styrene, naphthalene, ethylbenzene, and cyclohexane.<sup>48–51</sup> In addition, sMMO oxidizes various halogenated

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hydrocarbons, including pollutants such as trichloroethylene.<sup>10,52</sup> Crystallographic data indicate that substrates and products can occupy a chain of hydrophobic cavities, denoted cavities 1–3, in the  $\alpha$  subunit of MMOH (Figure 6A).<sup>53</sup>

**Image****Content**

**Figure 6.** Substrate binding by MMOH. (A) Three cavities and a pore region connect to the active site: cavity 1, green; cavity 2, wheat; cavity 3, light blue; hydrophilic pore, magenta. The iron ions are shown as orange spheres. (B) Structure of active site with bound 2-bromoethanol (PDB accession code 1XVG). The carbon atoms of 2-bromoethanol are shown in green and the bromine atom is shown in magenta.

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Cavities 2 and 3 are linked, but are only connected with cavity 1, which is next to the diiron active site, when MMOB is bound to MMOH.<sup>54,55</sup> A number of crystal structures with bound xenon, halogenated alkanes, and product alcohols have been obtained (Table 1). The substrate analogues dibromomethane, iodoethane, and xenon bind in cavities 2 and 3.<sup>56</sup> Along with methanol and ethanol, the product analogues 2-bromoethanol, 3-chloropropanol, and 3-bromobutanol bind in cavities 1–3 and can bridge the diiron center, replacing a bound water molecule (Figure 6B).<sup>54,57</sup> These structures are consistent with optical, resonance Raman, magnetic circular dichroism, electron nuclear double resonance (ENDOR), and EPR spectroscopic data showing that substrates and products can affect the diiron site.<sup>58–62</sup> However, recent time-resolved resonance Raman data indicate that the product alcohol does not bind directly to the diiron center.<sup>63</sup> Access to the active site cavity (cavity 1) from cavities 2 and 3 is mediated by conformational changes in two residues, Leu110 and Phe188, which form a gate.<sup>53,54</sup> The functional relevance of this chain of cavities to the catalytic cycle remains controversial, however. Finally, an  $\sim 10\text{ \AA}$  long hydrophilic pathway called the pore region is present and may

function in alcohol and water release as well as play a role in proton transfer.<sup>54,55,64</sup>

**Particulate MMO.** Because of difficulties isolating stable and active purified samples of pMMO, the enzyme kinetics have not been investigated thoroughly. Assays are further complicated by the use of different reductants, including NADH and duroquinol for isolated membranes, duroquinol for purified pMMO,<sup>8</sup> and formate for whole cells.<sup>48</sup> For isolated *M. capsulatus* (Bath) membranes, of which pMMO is the main component (such preparations are referred to as membrane-bound pMMO and can sometimes be crystallized upon solubilization without further purification<sup>25</sup>), methane oxidation activities are in the range of 25–130 nmol min<sup>-1</sup> mg<sup>-1</sup>, depending on the preparation and reductant.<sup>26</sup> Assuming that these samples are 80% pMMO, a turnover number in the range of 3–16 min<sup>-1</sup> can be estimated (Table 2). For propylene, membrane-bound pMMO activities are 16–123 nmol min<sup>-1</sup> mg<sup>-1</sup>,<sup>12,65</sup> giving an estimated turnover number of 2–15 min<sup>-1</sup>. For purified *M. capsulatus* (Bath) pMMO, specific activities of 17–160 nmol min<sup>-1</sup> mg<sup>-1</sup> for propylene have been reported,<sup>10</sup> corresponding to turnover numbers of 0.17–16 min<sup>-1</sup>. Lower activities of 2–23 nmol min<sup>-1</sup> mg<sup>-1</sup> have been obtained for membrane-bound pMMO samples from *M. trichosporium* OB3b, *Methylocystis* sp. str. M, and *Methylocystis* sp. str. Rockwell, with very little activity after solubilization and purification.<sup>9,26</sup> The whole cell *M. trichosporium* OB3b pMMO *V*<sub>max</sub> is 82–300 nmol min<sup>-1</sup> mg<sup>-1</sup> total protein (measured by methane consumption),<sup>66</sup> which corresponds to a turnover number of 30–150 min<sup>-1</sup>, assuming 20% of the total protein is pMMO.<sup>67</sup> Similar *V*<sub>max</sub> values were measured by propylene epoxidation for whole cells of *M. capsulatus* (Bath) pMMO.<sup>12,67,68</sup> Thus, preserving pMMO activity upon removal from the membranes remains an important issue. Apparent pMMO *K*<sub>m</sub> values for methane have been reported in the range of 1–62 μM.<sup>66,69,70</sup> Notably, some methanotrophs, including *Methylocystis* sp. strain SC2, can produce a high affinity pMMO with a much lower apparent *K*<sub>m</sub> for methane.<sup>70</sup> The molecular basis for this enhanced methane affinity is not known.

pMMO has a much narrower substrate range than sMMO and can oxidize only C1–C5 *n*-alkanes and terminal alkenes to 2-alcohols and 1,2-epoxides.<sup>48,71–73</sup> Analysis of the pMMO structures reveals a cavity at the proposed dicopper active site as well as a second cavity extending to the zinc binding site.<sup>19,74</sup> However, the functional relevance of these cavities remains unclear due to the moderate resolution of the pMMO structures and the lack of structures in the presence of substrates and products or analogues thereof. A hydrophobic pocket the size of pentane in the pmoA subunit has also been identified computationally<sup>75</sup> but is not near any of crystallographically observed metal binding sites. The narrow substrate specificity of pMMO and its ability to selectively orient substrates to produce (R)-alcohols<sup>72,76</sup> suggests that a fairly restricted cavity must be present at the active site. Site-directed mutagenesis studies of the *Mycobacterium* NBB4 HMO suggest that residue Ala 151, which is equivalent to pmoC Asp166 in *M. capsulatus* (Bath) pMMO, is important for determining substrate specificity.<sup>40</sup> This residue lies at the interface of pmoB and pmoC, about 8 Å from the dicopper center, and could modulate the sizes of the aforementioned cavities. Notably, AMO has a much broader substrate specificity,<sup>6</sup> but there are no crystal structures of AMO, and the molecular basis for this difference remains unclear.

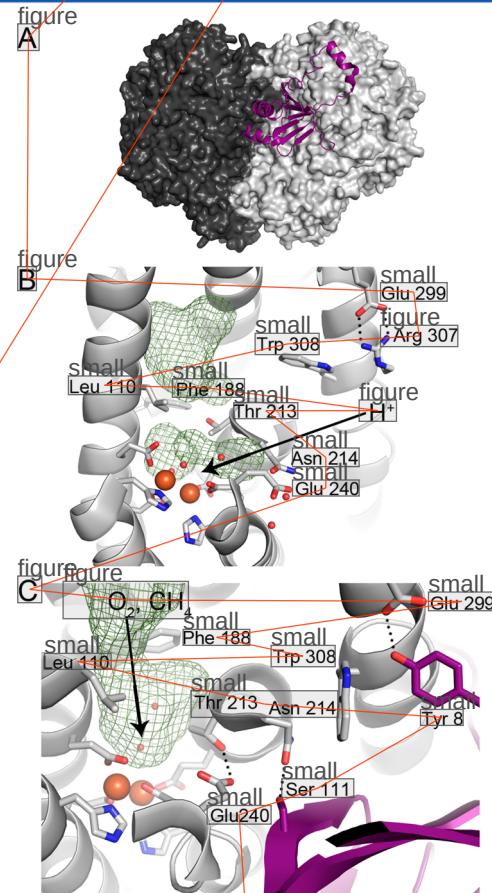
## Headline

## PROTEIN–PROTEIN INTERACTIONS

## Content

**Soluble MMO.** MMOH interacts with MMOB and MMOR. MMOH is essential for efficient hydrocarbon oxidation by MMOH. Binding of MMOB increases the reaction rate of reduced MMOH with O<sub>2</sub> by 1000-fold,<sup>77,78</sup> lowers the redox potential of the diiron center,<sup>79</sup> changes the regioselectivity of substrate oxidation,<sup>80</sup> and affects the structure of the diiron center.<sup>59,81,82</sup> The 2.9 Å resolution structure of the *M. capsulatus* (Bath) MMOH–2MMOB complex reveals that MMOB binds to the canyon region at the dimer interface of MMOH, leading to the ordering of the N-terminal 35 residues of MMOB, which are not observed in the NMR structures,<sup>83,84</sup> into a ring-like fold (Figure 7A).<sup>55</sup> MMOB binding also causes

## Image



## ContentDescription

Figure 7. Structural changes in MMOH upon complexation with MMOB. (A) MMOB (purple cartoon) binds in the canyon region formed by the two MMOH protomers (black and gray surfaces) (PDB accession code 4GAM). (B) Active site region in uncomplexed MMOH (PDB accession code 1MTY). The hydrophilic pore is open to solvent, and cavities 1 and 2 are separated by Leu110 and Phe188. (C) Active site region in MMOH–MMOB complex (PDB accession code 4GAM). The pore is closed, and conformational changes in Phe188 link cavities 1 and 2, allowing substrate access to the active site. Cavities are shown as green mesh.

## Content

conformational changes in MMOH residue Phe188 of the α subunit, opening up a pathway from the surface to the active site. In addition, Thr213 of the MMOH α subunit rotates inward toward the active site, forming new hydrogen bonds with Glu240 and closing the hydrophilic pore (Figure 7B,C). This residue has been proposed to play a role in proton delivery. Finally, MMOB binding induces small changes in

some of the ligands, including a shift of Glu243 to a coordination more similar to that in reduced MMOH.<sup>55</sup>

MMOR transfers two electrons from NADH one at a time to the diiron site in MMOH via its FAD and [2Fe-2S] cofactors.<sup>85,86</sup> There is no crystal structure of the MMOH-MMOR complex, but NMR titrations indicate that residues 24–34 in  $\alpha$  helix 1 and residues 67–69 in  $\alpha$  helix 2 interact with MMOH.<sup>87</sup> Chemical cross-linking studies have suggested that MMOR interacts with both the  $\alpha^{88}$  and  $\beta^{81}$  subunits of MMOH, and recent work using hydrogen–deuterium exchange coupled to mass spectrometry (HDX-MS) and computational docking suggests that the ferredoxin domain of MMOR binds to MMOH at the same location where MMOB binds. At this binding site, the MMOR [2Fe-2S] cluster is positioned 14 Å away from the diiron active site and does indeed interact with both subunits. Consistent with these findings, MMOB inhibits MMOR binding as well as electron transfer.<sup>89</sup> However, it is also possible that MMOB inhibits MMOR binding by forming a complex with MMOR.<sup>81</sup>

Recent fluorescence anisotropy experiments suggest that MMOB has a higher affinity for MMOH in the reduced form ( $K_d = 0.17 \mu\text{M}$ ) than in the oxidized form ( $K_d = 0.55 \mu\text{M}$ ),<sup>90</sup> an observation that is difficult to reconcile with the lowered redox potential of MMOH in the presence of MMOB.<sup>79,91</sup> Double electron–electron resonance (DEER) spectroscopic data further suggest that the conformation of the N-terminal tail of MMOB is more rigid when MMOH is reduced.<sup>90</sup> The key role of the MMOB N-terminus is underscored by data showing that mutants of MMOB disrupting the N-terminus display significantly reduced activity.<sup>55,92</sup> In one functional model, MMOR is proposed to replace MMOB, transfer electrons to reduce the active site, and then MMOB is proposed to rebind to the reduced form, displacing MMOR, and initiating O<sub>2</sub> activation and methane oxidation.<sup>89</sup> In the absence of experimental evidence for MMOR displacing MMOB from the oxidized MMOH, this scenario remains speculative.

**Particulate MMO.** Protein–protein interactions are not required for pMMO activity in vitro, but protein–protein complexes involving pMMO have been suggested. Besides the interaction with the mystery helix observed crystallographically (Figure 3),<sup>24–26</sup> pMMO has been proposed to interact with the next enzyme in the methane oxidation metabolic pathway, methanol dehydrogenase (MDH).<sup>93</sup> MDH is a periplasmic enzyme that oxidizes methanol to formaldehyde and is composed of a 63 kDa  $\alpha$  subunit, which contains a pyrroloquinoline quinone (PQQ)/calcium ion cofactor, and a 8 kDa  $\beta$  subunit arranged as an  $\alpha_2\beta_2$  dimer.<sup>94,95</sup> The *M. capsulatus* (Bath) MDH can also form higher order heterogeneous oligomers.<sup>95</sup> A putative pMMO–MDH supercomplex was isolated, and its structure was determined to 16 Å resolution by electron microscopy. This structure was interpreted as three  $\alpha\beta$  MDH monomers capping the periplasmic pmoB soluble domains of the pMMO  $\alpha_3\beta_3\gamma_3$  trimer.<sup>96</sup> This pMMO–MDH complex exhibited moderately higher propylene epoxidation activity than pMMO alone,<sup>96,97</sup> but its methane oxidation activity was not reported. Recent biolayer interferometry studies of MDH and pMMO from *M. capsulatus* (Bath) reveal a specific protein–protein interaction.<sup>95</sup> An interaction was also detected between MDH and the spmoB recombinant protein, indicating that the periplasmic regions of pmoB are involved.<sup>95</sup> These data are consistent with reports that MDH is associated with the membranes in various methanotrophs.<sup>98–100</sup> The observed complex is likely dynamic

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since a stable complex cannot be isolated by size exclusion chromatography, and its formation may depend on the presence of the membrane or other proteins. One proposal is that the lysine-rich  $\beta$  subunit of MDH, of which the function is not known, interacts with pMMO and/or negatively charged phospholipid headgroups at the periplasmic membrane,<sup>95</sup> thus associating with pMMO in a fashion that would be difficult to reconstitute in vitro. Such a model would not apply to methanotrophs that utilize a subset of MDH enzymes lacking the  $\beta$  subunit,<sup>101</sup> but these methanotrophs, which include the Verrucomicrobia,<sup>102</sup> also have different types of membranes.<sup>36</sup>

A pMMO/MDH supercomplex would provide a direct pathway for methanol produced at the proposed dicopper active site in the pMMO pmoB subunit to the active site of MDH where it is oxidized to formaldehyde. In addition, such a complex could directly provide reducing equivalents to pMMO through the cytochrome c<sub>1</sub>, the electron acceptor of MDH.<sup>96,97,103–105</sup> Another and perhaps more widely accepted electron source for pMMO involves ubiquinol generated by a type 2 NADH:quinone oxidoreductase (NDH-2).<sup>12,106,107</sup> In support of this pathway, NADH is an effective reductant for membrane-bound pMMO, but duroquinol is the only reductant that is effective after detergent solubilization.<sup>71,108</sup> Duroquinol may mimic the function and redox potential of endogenous quinols,<sup>108</sup> and it is known that methanotrophs produce a range of ubiquinones, but not menaquinone.<sup>109,110</sup> Moreover, addition of NADH and NDH-2 to purified pMMO samples leads to a modest increase in propylene epoxidation activity.<sup>12</sup> NDH-2 and pMMO can be copurified,<sup>12</sup> but there is no evidence that they form a specific protein–protein complex. Finally, in *M. capsulatus* (Bath), a cytoplasmic hemerythrin protein increases the activity of pMMO, suggesting that hemerythrin facilitates oxygen delivery to pMMO.<sup>111</sup> However, there is no evidence for a protein–protein interaction between pMMO and this hemerythrin.

## heading

## MECHANISM

**Soluble MMO.** The details of how sMMO activates dioxygen have been probed extensively by single turnover kinetic studies, resulting in the identification of transient oxygen intermediates, the kinetic and spectroscopic properties of which are detailed elsewhere.<sup>14,19,112</sup> First, the reduced diiron(II) MMOH reacts with O<sub>2</sub> in the presence of 2 equiv of MMOB to form a species O in which O<sub>2</sub> is bound, but not directly to the diiron cluster.<sup>77</sup> Intermediate O is then converted to peroxy intermediates P\* and P. For *M. capsulatus* (Bath) MMOH, P\* is observed to be a diiron(III) site that differs from the subsequent intermediate P only in ligand arrangement.<sup>113</sup> However, recent studies of *M. trichosporium* OB3b MMOH utilizing a variant of MMOB that slows conversion of P\* to P indicate that P\* contains a diiron(II) site similar in structure to the diiron(II) site in reduced MMOH. In this system, P\* is proposed to form by displacement of the bridging H<sub>2</sub>O ligand by O<sub>2</sub> without electron transfer from the diiron(II) center.<sup>114</sup> Thus, it remains unclear whether the two intermediates denoted P\* are the same. The second peroxy intermediate, P, is a diiron(III) species, and spectroscopic comparisons as well as computational work suggest that a  $\mu$ -1,2-peroxy bridge is likely.<sup>14,115</sup> Formation of P is pH-dependent and exhibits a kinetic solvent isotope effect in D<sub>2</sub>O, indicating that proton transfer plays a role.<sup>113,116</sup> It is also possible to bypass the requirement for MMOB and MMOR in these steps by reacting the oxidized MMOH with H<sub>2</sub>O<sub>2</sub> as source of both oxygen and

electrons. However, very high concentrations of  $\text{H}_2\text{O}_2$  are required, with an apparent  $K_m$  value of  $\sim 66 \text{ mM}$ .<sup>117,118</sup>

Intermediate P is then converted to Q in a proton-dependent step<sup>113,116</sup> that results in O–O bond breaking. Intermediate Q is the species that reacts with methane to break the C–H bond. On the basis of Mössbauer spectroscopic data indicating the presence of two Fe<sup>IV</sup> ions in similar coordination environments and EXAFS data best fit with an Fe–Fe distance of 2.46 Å, Q was proposed to adopt a di( $\mu$ -oxo) diamond core structure.<sup>119,120</sup> Difficulty reproducing this distance computationally<sup>121,122</sup> combined with the observation that a model complex with an open core terminal Fe<sup>IV</sup>=O unit can oxidize C–H bonds rapidly,<sup>123</sup> although not those in methane, suggested Q might have an alternative structure. However, a recent time-resolved resonance Raman study reveals a vibration from Q that best matches vibrations observed for the diamond core in model compounds. Additional experiments with mixed isotopes of O<sub>2</sub> further indicate that Q forms via homolytic cleavage of the O–O bond.<sup>63,124</sup> Once Q is formed, there are several mechanistic possibilities for C–H activation, including both concerted and radical or cationic processes.<sup>19</sup> A range of approaches, such as the use of chiral alkane and radical clock substrates, the measurement of kinetic isotope effects, and computational analysis, have been used to investigate these options.<sup>125</sup> After methane oxidation, Q is converted to the product complex T, which is proposed to have a mono- $\mu$ -oxo bridged structure.<sup>63</sup>

**Particulate MMO.** Mechanistic studies of pMMO are far behind those of sMMO, due to ongoing difficulties isolating highly active purified enzyme and uncertainty as to the atomic details of the active site (vide supra).<sup>9</sup> Density functional theory (DFT) and quantum mechanics/molecular mechanics (QM/MM) studies suggest that a monocupper Cu<sup>III</sup>-oxo species or a di( $\mu$ -oxo)Cu<sup>II</sup>Cu<sup>III</sup> species could oxidize methane and that formation of the dicopper species is more favorable.<sup>126</sup> Calculations using a dicopper active site model suggest that either a  $\mu\text{-}\eta^1\text{:}\eta^2$ -peroxy Cu<sup>I</sup>Cu<sup>II</sup> or  $\mu\text{-}\eta^2\text{:}\eta^2$ -peroxy Cu<sup>II</sup>Cu<sup>II</sup> species is formed first and then is transformed into the more reactive di( $\mu$ -oxo)Cu<sup>II</sup>Cu<sup>III</sup> species after O–O bond cleavage.<sup>126,127</sup> In later studies, a ( $\mu$ -oxo)( $\mu$ -hydroxo)Cu<sup>II</sup>Cu<sup>III</sup> center was proposed to be the active species and was calculated to be more reactive toward methane than a di( $\mu$ -oxo)Cu<sup>II</sup>Cu<sup>III</sup> species. Formation of this ( $\mu$ -oxo)( $\mu$ -hydroxo)Cu<sup>II</sup>Cu<sup>III</sup> species is proposed to be facilitated by proton transfer involving residues Tyr374 and Glu35 (M. capsulatus (Bath) numbering; Tyr374 is not structurally conserved in other pMMOs) in the second coordination sphere of the dicopper center.<sup>128</sup> It is important to note that these intermediates have been calculated based on the moderate resolution crystal structures of pMMO and have not been detected experimentally. Related to these ideas are recent studies of the zeolite Cu-ZSM-5, which can oxidize methane to methanol. Spectroscopic studies and DFT calculations show that the zeolite active species is a mono( $\mu$ -oxo)Cu<sup>II</sup>Cu<sup>II</sup> core formed from a  $\mu\text{-}\eta^2\text{:}\eta^2$ -peroxyCu<sup>II</sup>Cu<sup>II</sup> intermediate.<sup>129,130</sup> Although pMMO and the zeolite are fundamentally different, this work does suggest that it may be feasible to oxidize methane in an enzyme with a dicopper site.

One putative oxygen intermediate has been detected experimentally for pMMO. When reduced pMMO is reacted with H<sub>2</sub>O<sub>2</sub> or reduced spmoB is reacted with O<sub>2</sub> or H<sub>2</sub>O<sub>2</sub>, a new species with an absorbance feature at 345 nm is formed.<sup>131</sup> The energy and intensity of this feature are consistent with a  $\mu\text{-}\eta^2\text{:}\eta^2$ -peroxy Cu<sup>II</sup>Cu<sup>II</sup> species like those formed in oxy-

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hemocyanin and oxytyrosinase,<sup>132</sup> a met Cu<sup>II</sup>Cu<sup>II</sup> species like that in methemocyanin,<sup>133,134</sup> or a hydroxo-bridged Cu<sup>II</sup>Cu<sup>II</sup> center like that in the type 3 copper centers of multicopper oxidases.<sup>135</sup> Because of the instability of spmoB and heme contaminants present in pMMO, further studies of this species by resonance Raman spectroscopy have not been conducted. Importantly, this intermediate does not form in variants of spmoB in which the ligands to the dicopper center have been mutated, consistent with O<sub>2</sub> binding at the proposed active site. In addition, this species disappears in the presence of methane,<sup>131</sup> and some methanol product is detected, indicating that it may be a relevant intermediate during methane oxidation. The identity of the actual species that reacts with methane, the pMMO equivalent to Q, remains unclear. Several studies using chiral alkanes suggest that C–H bond activation by this unknown species proceeds via a concerted, nonradical mechanism.<sup>136</sup>

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## CONCLUSIONS

## Content

The two methane-oxidizing enzymes in nature, sMMO and pMMO, are completely different. The sMMO hydroxylase component, MMOH, is a soluble  $\alpha_2\beta_2\gamma_2$  dimer, whereas pMMO is an  $\alpha_3\beta_3\gamma_3$  trimer with both membrane-bound and periplasmic regions. The active site of sMMO is a well-characterized diiron center, but the atomic structure of the pMMO copper active site remains unclear. Similarly, the structural basis for substrate and product access to the sMMO active site has been partially elucidated but is not well-defined for pMMO. Both MMOs oxidize a range of hydrocarbon substrates, but pMMO has a narrower substrate specificity. sMMO exhibits higher turnover numbers than pMMO both in cells and in isolated enzyme preparations, but neither enzyme achieves levels desirable for biological gas-to-liquid conversion processes. The delivery of electrons to the sMMO active site in MMOH and the roles of MMOB and MMOR are reasonably well understood on the molecular level, although some aspects remain controversial. By contrast, the source of electrons for pMMO is still under debate, and possible protein interaction partners such as MDH require further investigation. The catalytic cycle of sMMO has been mapped out, and the intermediate that breaks the methane C–H bond, Q, has been characterized. Although mechanistic possibilities for pMMO have been considered, its mechanism cannot be further elucidated until lingering questions about the active site are resolved. Despite all these differences, the two systems share the unique ability to oxidize methane, and it is remarkable that nature has evolved two distinct large enzyme complexes for oxidation of a small gaseous substrate. Further study of how each MMO orchestrates the delivery of methane, oxygen, protons, and electrons to the active site, and how the O–O and C–H bonds are broken is essential to understanding these enzymes, their roles in the environment, and their potential biotechnological applications.

## authors

## AUTHOR INFORMATION

## authors

## Corresponding Author

\*E-mail: amyr@northwestern.edu.

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Notes

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**ABBREVIATIONS**

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AMO, ammonia monooxygenase; ENDOR, electron nuclear double resonance; EPR, electron paramagnetic resonance; EXAFS, X-ray absorption fine structure; HMO, hydrocarbon monooxygenase; MDH, methanol dehydrogenase; MMO, methane monooxygenase; MMOB, sMMO regulatory protein; MMOR, sMMO reductase; MMOH, sMMO hydroxylase; NDH-2, type 2 NADH:quinone oxidoreductase; pMMO, particulate methane monooxygenase; sMMO, soluble methane monooxygenase; spinoB, soluble fragment of the pMMO pmoB subunit

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