Crystal structure of the hydroxylase component of methane monooxygenase from *Methylosinus* trichosporium OB3b

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Abstract

Methane monooxygenase (MMO), found in aerobic methanotrophic bacteria, catalyzes the O₂-dependent conversion of methane to methanol. The soluble form of the enzyme (sMMO) consists of three components: a reductase, a regulatory "B" component (MMOB), and a hydroxylase component (MMOH), which contains a hydroxo-bridged dinuclear iron cluster. Two genera of methanotrophs, termed Type X and Type II, which differ markedly in cellular and metabolic characteristics, are known to produce the sMMO. The structure of MMOH from the Type X methanotroph Methylococcus capsulatus Bath (MMO Bath) has been reported recently. Two different structures were found for the essential diiron cluster, depending upon the temperature at which the diffraction data were collected. In order to extend the structural studies to the Type II methanotrophs and to determine whether one of the two known MMOH structures is generally applicable to the MMOH family, we have determined the crystal structure of the MMOH from Type II Methylosinus trichosporium OB3b (MMO OB3b) in two crystal forms to 2.0 Å and 2.7 Å resolution, respectively, both determined at 18 °C. The crystal forms differ in that MMOB was present during crystallization of the second form. Both crystal forms, however, yielded very similar results for the structure of the MMOH. Most of the major structural features of the MMOH Bath were also maintained with high fidelity. The two irons of the active site cluster of MMOH OB3b are bridged by two OH (or one OH and one H₂O), as well as both carboxylate oxygens of Glu α 144. This bis- μ hydroxo-bridged "diamond core" structure, with a short Fe-Fe distance of 2.99 Å, is unique for the resting state of proteins containing analogous diiron clusters, and is very similar to the structure reported for the cluster from flash frozen (-160 °C) crystals of MMOH Bath, suggesting a common active site structure for the soluble MMOHs. The high-resolution structure of MMOH OB3b indicates 26 consecutive amino acid sequence differences in the β chain when compared to the previously reported sequence inferred from the cloned gene. Fifteen additional sequence differences distributed randomly over the three chains were also observed, including D α 209E, a ligand of one of the irons.

Keywords: dinuclear iron cluster; methane oxidation; methanotroph; oxygen activation; X-ray crystallography

Methanotrophic bacteria utilize methane produced from the metabolism of anaerobic bacteria and other natural sources to satisfy their total carbon and energy requirements (Dalton, 1980; Anthony, 1982). The first and most difficult step of the methane oxidation pathway, in which the stable C-H bond of methane is broken and an oxygen atom inserted to form methanol, is catalyzed

by the enzyme methane monooxygenase (MMO) (Dalton, 1980; Feig & Lippard, 1994; Lipscomb, 1994; Wallar & Lipscomb, 1996):

 $CH_4 + O_2 + NAD(P)H + H^+ \rightarrow CH_3OH + H_2O + NAD(P)^+.$

MMO also catalyzes the adventitious oxidation of a large number of other hydrocarbons up to about C_8 in size (Colby et al., 1977; Dalton, 1980; Higgins et al., 1980; Green & Dalton, 1989; Tsien et al., 1989; Fox et al., 1990b; Andersson et al., 1991; Rataj et al., 1991). MMO exists in both a soluble (sMMO) and a particulate (pMMO) form (Dalton, 1980; Anthony, 1982). sMMO has been purified and characterized from the Type X methanotroph, Methylococcus capsulatus Bath (MMO Bath) (Dalton, 1980, 1991; Woodland & Dalton, 1984; Pilkington & Dalton, 1990), and several Type II methanotrophs including, Methylosinus trichosporium OB3b

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(MMO OB3b) (Fox et al., 1989, 1990a), and *Methylobacterium* sp. CRL-26 (Patel & Savas, 1987). The Type X and Type II methanotrophs differ dramatically in cellular morphology, many metabolic pathways, and response to environmental stress (Anthony, 1982), but their MMOs and other methane oxidation pathway enzymes appear to be very similar.

The MMO operons from M. trichosporium OB3b and M. capsulatus Bath have been cloned and sequenced (Stainthorpe et al., 1989, 1990; Cardy et al., 1991a, 1991b). These studies and direct protein isolation and characterization have shown that MMO is composed of three protein components, each of which is required for efficient substrate hydroxylation coupled to NADH oxidation: a 245-kDa hydroxylase (MMOH), a 40-kDa reductase (MMOR), and a 15-kDa protein called component B (MMOB). The hydroxylase has a dimeric protomer structure, with each protomer containing three subunits (α, β, γ) (Rosenzweig et al., 1993; Froland et al., 1994) and a hydroxo-bridged dinuclear iron cluster (Fox et al., 1988, 1993; DeWitt et al., 1991; DeRose et al., 1993; Thomann et al., 1993). MMOH and its diiron clusters are essential for catalysis by MMO (Fox et al., 1989). The cluster is contained in the α subunit adjacent to an internal hydrophobic pocket that forms the active site (Rosenzweig et al., 1993). The genetic studies indicate a 66% sequence identity between the hydroxylase subunits of the two species that have been cloned and sequenced (Stainthorpe et al., 1989, 1990; Cardy et al., 1991a, 1991b).

The other two components of MMO also serve important functions. In addition to transferring electrons to MMOH, MMOR forms a specific complex with the β subunit of MMOH (Fox et al., 1991), which has the effect of adjusting the relative redox potentials of the two cluster irons so that two-electron transfer is favored (Paulsen et al., 1994; Y. Liu, J. Nesheim, K.E. Paulsen, M.T. Stankovich, & J.D. Lipscomb, manuscript submitted). Component B has been shown to bind specifically to the α subunit of MMOH (Fox et al., 1991). This complex has profound effects on the rate of certain reactions in the catalytic cycle, the redox potential of the diiron cluster, the structure of MMOH, and the product distribution and yield of the overall reaction (Fox et al., 1991; Froland et al., 1992; Liu et al., 1995b, 1995c).

All steps of the catalytic cycle of sMMO appear to involve the diiron cluster (Fig. 1) (Fox et al., 1989; Lee et al., 1993b). The first step in the catalytic cycle is the reduction of the diiron cluster to the diferrous state (Fox et al., 1989). Next, O₂ binds to the cluster in two steps, forming a peroxo species termed compound P (Lee et al., 1993b; Liu et al., 1994, 1995a, 1995c). This species then converts spontaneously to a high valent iron-oxo species (compound Q) (Lee et al., 1993a, 1993b) in which both irons appear to be in the Fe(IV) state. This species is electronically and functionally analogous to the proposed mononuclear Fe(IV) oxenoid π cation radical species (compound I) produced by cytochrome P-450 and horseradish peroxidase during catalysis (Ortiz de Montellano, 1985; McMurry & Groves, 1986). At this stage, the highly oxidized diiron cluster reacts with substrate to form the hydroxylated product and the resting diferric state of MMOH after passing through other intermediates that have been characterized partially (Priestley et al., 1992; Lee et al., 1993b; Nesheim & Lipscomb, 1996).

The crystal structure of the hydroxylase component from the Type X methanotroph *M. capsulatus* Bath has been reported in the oxidized state at 4 °C and -160 °C (Rosenzweig et al., 1993, 1995) as well as the reduced state at -160 °C (Rosenzweig et al., 1995). Interestingly, the structure of the dinuclear iron cluster was found to be different in each of the conditions and redox states. In par-



Proposed Catalytic Cycle for MMO



P-450 Activated Species

Fig. 1. Proposed catalytic cycle of methane monooxygenase based on a radical recombination process for substrate hydroxylation. The cytochrome P-450 activated species is the proposed mononuclear Fe(IV) oxenoid π cation radical species equivalent to compound Q. Oxygen atoms derived from O₂ are filled.

ticular, the two structures of the cluster in the resting diferric enzyme differed significantly in that, at 4 °C, the two irons were bridged by Glu α 144, a hydroxide moiety, and an exogenous acetate to give an Fe-Fe distance of 3.4 Å, whereas the structure determined at -160 °C revealed Glu α 144, a hydroxide, and a water as bridging ligands to give an Fe-Fe distance of 3.1 Å. The latter structure is consistent with model compounds for the diiron clusters containing a so-called "diamond core" with two bridging oxygen ligands (see, for example, Zang et al., 1994; Que & Dong, 1996), as well as with EXAFS studies of MMOH OB3b that indicated a predominant Fe-Fe distance of about 3 Å (Shu et al., 1996a).

In order to determine whether one of the structures of diferric MMOH Bath is characteristic of MMOHs in general, and to extend structural studies to the Type II methanotrophs, we have determined the X-ray crystal structure of MMOH OB3b. Reported here are the 18 °C structures of MMOH OB3b to 2 Å resolution and of MMOH OB3b crystallized in the presence of MMOB to 2.7 Å resolution. We find that the overall structures of MMOHs from the Type X and Type II methanotrophs are very similar, and that the MMOH OB3b forms contain a diiron cluster with a short Fe-Fe distance [2.99 Å in crystal form 1 (MMOH alone) and 3.07 Å in crystal form 2 (MMOB present during crystallization)] and a diamond core oxygen bridge structure. This suggests that the diamond core structure is a common characteristic of MMOH from these widely divergent methanotrophs and may be an important feature for catalysis.

Results and discussion

Structure determination

The structure of MMOH OB3b in crystal form 1 was solved by molecular replacement using a monomer of MMOH Bath (Rosenzweig et al., 1993) as the search model and the program XPLOR (Brünger, 1992). Cycles of molecular dynamics refinement and Powell minimization were alternated with map examination, model building, and refinement, to produce the final model that has a crystallographic *R*-factor of 13.7% for data between 5 and 2 Å, with $F > 2\sigma$ (14.5% with no cut off). This model has 9,334 non-hydrogen atoms from 510 residues in the α chain, 383 residues in the β chain, 167 residues in the γ chain, 731 water molecules, two iron atoms, and two hydroxides. Residues $\alpha 1 - \alpha 16$, $\beta 1-\beta 10$, $\beta 393-\beta 394$, $\gamma 1$, and $\gamma 169$ were disordered and not included in the final model. Overall, the electron density map is exceptionally clear, except for the few disordered residues. Figure 2 shows two regions of $2F_o - F_c$ electron density that are typical in quality. Ramachandran plots for the three chains indicate that 93.5% of the residues have (ϕ, ψ) values in the "most favored" region, 6.4% in the "additional allowed" region, and one residue in the "disallowed region" as identified by PROCHECK (Laskowski et al., 1993). Ser γ 42 is the only residue in the disallowed region of the Ramachandran plot and is found in an exposed surface loop.

The structure of MMOH OB3b in crystal form 2 was solved using molecular replacement with a refined model of MMOH OB3b in crystal form 1 as a search model. Rigid-body minimization followed by molecular dynamics refinement reduced the R-factor to 19.5% for data between 8 and 2.8 Å, with $F > 2\sigma$. Examination of the electron density maps followed by Powell minimization produced the final model with a crystallographic R-factor of 15.2% for data between 5 and 2 Å, with $F > 2\sigma$ (17.8% with no cut off). This model has 8,605 non-hydrogen atoms from 510 residues in the α chain, 383 residues in the β chain, 167 residues in the γ chain, two water molecules, two iron atoms, and two hydroxides. Ramachandran plots for the three chains indicate that 90.5% of the residues have (ϕ, ψ) values in the "most favored" region, 9.1% in the "additional allowed" region, two residues in the generously allowed region, and one residue in the "disallowed region" as identified by PROCHECK (Laskowski et al., 1993). Again, Ser γ 42 is the only residue in the disallowed region of the Ramachandran plot.

A summary of the data collection, refinement, and geometric quality for both of the crystal forms of MMOH OB3b is given in the statistical parameters supplied in Table 1.

Overall structure

The sequence alignment of MMOH OB3b and MMOH Bath with the secondary structural elements labeled to be consistent with that of MMOH Bath is shown in Figure 3. MMOH is almost exclusively helical, with 19 helices in the α chain, 12 helices in the β chain, and 8 helices in the γ chain, accounting for 634 residues of the 1,060 residues in the protomer. There are only three short strands of β structure, accounting for 15 residues.

The three-dimensional structure of MMOH OB3b is shown in Figure 4. The crystallographic twofold, coincident with the crystallographic *a* axis and relating the two protomers, is vertical in the figure. The heart-shaped dimer is about 130 Å \times 90 Å \times 70 Å, with the α chains forming the auricles and the β chains forming the ventricles. Although not immediately obvious from the figure, the

 α and β chains share a similar fold. Two-hundred thirty-three pairs of $C_{\alpha}s$ can be superimposed with an RMS difference of 2.08 Å. This fold is also found in the R2 protein of ribonucleotide reductase (Nordlund et al., 1990). The interface between the α and β chains is extensive, burying 4,068 Å² of the solvent-exposed surface of each chain. Each γ chain is elongated and interacts with the α and β chains away from the symmetry axis.

The active site, which contains the diiron cluster, is located in the α chain between the four long helices, $\alpha B (\alpha 97 - \alpha 127)$, $\alpha C (\alpha 131 - \alpha 161)$, $\alpha E (\alpha 197 - \alpha 226)$, and $\alpha F (\alpha 231 - \alpha 257)$. The distance between the symmetry-related active sites is about 45 Å. A more detailed description of the active site and metal center will be given below.

Several of the helices ($\alpha 1$, αD , αE , αH , βB , βE , βF , and βG) are broken into smaller segments. Most of these distortions occur at one end, sometimes involving 3₁₀ or π helical loops. Three of these helices should be noted. Helix αH is interesting in that the central turn of this five-turn helix is a π loop. A basis for this distortion can be seen in the presence of a tryptophan on each turn of this 22-residue helix. These large residues lock side chains to particular locations, leading to this unusual distortion. Helices αE and βE are structurally homologous and are each composed of three shorter helices. The bowing of helix αE is in large part responsible for the formation of the active site cavity in addition to supplying one of the iron ligands (Glu $\alpha 209$).

Structures of form 1 and form 2 of MMOH OB3b are essentially identical. The molecules pack in their respective C222₁ space groups with a relative rotation angle of 13.2° around the crystallographic dyad. This rotation lengthens the *x* axis of form 2 from 264.49 Å to 293.38 Å and the *z* axis from 139.44 Å to 143.65 Å, and reduces the *y* axis from 71.19 Å to 64.01 Å. The volumes of the two unit cells differ by less than 2.5%. While superimposing only a monomer of form 1 on form 2, the other monomers of the dimer were related by a rotation of 1.2° and a translation of 0.14 Å. RMS difference due to superposition of the C_as of a monomer between the two forms of MMOH is 0.27 Å. However, there is no noticeable change observed at the interface of the form 2 monomers compared with form 1 of the MMOH OB3b.

Differences between observed and predicted amino acid sequence

During the course of the map examination and model building, several places were found where the electron density map did not agree with the reported amino acid sequence inferred from the gene sequence (Cardy et al., 1991a, 1991b). These sites are shown in Figure 3 and fall into two classes. First, there are 15 randomly distributed sequence differences where the electron density map indicated a sequence more similar to that of MMOH Bath. For eight of these differences (D α 209E, G α 226A, G α 357A, M β 255Y, I β 256D, D γ 109E, G γ 109A, and P γ 160R), the published sequence (Cardy et al., 1991b) would have been sterically possible, but strong (>5 σ) features in the $F_o - F_c$ maps and the definition of the $2F_{o} - F_{c}$ map left no doubt as to the discrepancy. Prominent in this group of differences is the change at $\alpha 209$ from aspartate to glutamate (see Fig. 2A,C) because this residue is a ligand of the diiron cluster. For seven of the differences, not only were the maps clear, but in three cases (R α 95G, insertion of P α 329, and R γ 88G) changes were required because the reported sequence was sterically impossible, and in four cases (R α 37W, T α 210A, I α 225S, and V α 331S), the reported residues were disfavored due to short steric contacts or the burying of unpaired charges.









Table 1. Standard crystallographic data

	Form 1	Form 2
Space group	C2221	C2221
Unit cell	a = 264.49 Å	a = 293.38 Å
	b = 71.19 Å	b = 64.01 Å
	c = 139.44 Å	c = 143.65 Å
Data collection		
Number of crystals	9	1
Number of reflections	391,667	177,040
Number of unique reflections	87,489	33,478
Completeness	96.8%	89.3%
R_{merge} of intensities ^a	0.075	0.094
Data used in final refinement		
Number of reflections	75,570	22,264
Cutoffs	5–2 Å, $F > 2\sigma$	5–2.7 Å, $F > 2\sigma$
Final R _{factor} ^b	0.137	0.152
Final R_{factor}^{b} with $F > 0\sigma$	0.145	0.178
Model		
Number of protein atoms	8,599	8,599
Number of water molecules	731	2
Number of iron atom	2	2
Number of hydroxides	2	2
$\langle \Delta r \rangle$ from Luzzati plot	0.15 Å	0.20 Å
Stereochemistry		
RMS nonideal bond lengths	0.016 Å	0.013 Å
RMS nonideal bond angles	1.65°	1.61°
RMS dihedral angle	21.14°	21.39°
RMS improper angles	1.58°	1.54°
Themal parameters		
$\langle B \rangle$ for protein atoms	19.2 Å ²	18.5 Å ²
$\langle B \rangle$ for water molecules	37.1 Å ²	20.4 Å ²
$\langle B \rangle$ for iron and hydroxides	15.5 Å ²	27.6 Å ²

$${}^{a}R_{merge} = \frac{\sum_{hkl}\sum_{i}|I_{i}(hkl) - \langle I(hkl)\rangle|}{\sum_{hkl}\sum_{i}I_{i}(hkl)}.$$

$${}^{b}R_{factor} = \frac{\sum_{hkl}|F_{obs}(hkl) - F_{calc}(hkl)|}{\sum_{hkl}F_{obs}(hkl)},$$

The second class of observed differences in the amino acid sequence is found near the carboxyl terminus of the β chain. This group of 26 consecutive differences is located at the top edge of the β chain as it is displayed in Figure 4. An examination of the published gene sequence corresponding to this region for MMOH OB3b (Cardy et al., 1991b) as shown in Figure 5 indicates that the observed amino acid sequence would be obtained if the base pair changes indicated in the figure occurred either through mutation of the strain or through sequencing errors. Figure 2B shows the exN. Elango et al.

a i	CH	AI	IN							
			10	20	30	W 40	50	60	70	
OB3 Bat	th th	1	MATSLATKAA MALSTATKAA	TDALEVNRAP TDALASNRAP	VOVEPORVER TEVNAQUEVER	WLQSPNRDFK WLQSPNRDFK	ENRTEYPTKY IMRTEYATKY	HMANETREOF IMANETREOF	KUIAKEYARM KLIAKEYARM	70 70
OBS	b	i.	EARDEROFO	TLLDGLTRLO	AGNEVHPRWG	ETMEVISHEL	EVGEYNALAA	SAMLADSATA	AEQENGYLAQ	140
Bat	h	I.	EAVEDERQFO	SLOVALTRLN	AGVEVEREN	ETHRVVSNFL	EVGEYNAIAA 038	TOMENDEAQA	AEQENGYLAQ	140
083	b	1	VLDEIRBTBO	CAFINRYYSE	HYRDPAGHND	ARRTRAIGPL	WEGMERVERD	GFISRDAVEC	SVHLQLVGDT	210
BAT	4	1	VLDEIRHTHQ	CATVBYFFAK	NUCOPAGIND	ARRTRTIOPL	WILCOMORRY IFILD	GFISGDAVEC	SINLQLVGUA	210
on	ь	1	CETHFLIVAY	TEWAIGNODE	ITPTVPLSVE	TDELREMANG	YOTVVSIAND	PASARFINTD	INNAPWTOOK	280
Bat	h	1	CETNELIVAY	TEWAAANGDE	ITPTVFLSIE	TDELREMANO	YOTVVBIAND	PASAEYLNTD	LINNAFWTQQK	280
011	ib.	1	YFTPVLOYLE	EYGSRFRVEP	WVKTWIGWVE	EDWOGIWIOR	LGRYGVES-R	VLEDAREDAY	WAHHDLALAA	350
Bat	th	1	YFTPVLORILE		WVETWDRWVY	EDWGGIWIGR	LOKYGVESPR	SLEDARQDAY $\leftarrow \alpha 4 \rightarrow$	WARHDLYLLA	350
OH:	1b		YAMWPLAFAR	LALPDEEDQA	WFEANYPOWA	DHYGEL FREW	FELOYEDPES	GFIPYQWLLA	NGHOVYIDRY	420
Bat	h	1	YALWPTOFFR	LALFDOREME (WFEANYPOWI	DHYOKIYEEW	RARGCEDPSS	OFIFINWEIE	NUMPITIDRY	420
OB:	tb	1	SQUPPIPSLA	KOTOSLAVHE	FNGERBLITD	DWGEROWLIE	PERYECHNVF	EQYEGRELSE	VIAEGHOVRS	490
Bat	h	1	BQVPFCPSIA	CIANTLEVHE CON	YNGEHHTFAD	QWGERMWLAE	PERYECQNIF	EQTEGRELSE	VIAELHGLRS X11→	490
OB:	lb	1	DORTLIAOPE	TRODULWTLE	DIRIGOCVPP	DPLAKE				
Bat	th	1	DOKTLIAQPE	VRODELNTLD	DIKRLNCVFE α12→	NPVRAF				
β	ĊH	A	IN							
OB.	3b	1	MEQPOSIQVI	FRGLTDPERA	ATTAAAVPDH	ALDTORKYHY	FIQPRWRPLS	EYEQLICTAQ	PNPDWIAGGL	70
Bat	th	3	MSHLGES	PROLTDPENA	AVILEALPEA	PLOSINIUGY	PUTPRWKRLT	EYEALTVYAQ ← D2 →	PHADWIAGGL	67
OB.	Лb	1	DWODWTQKFT	I GORPSWONES	TELETTOWYR	HRDPARRWII	PYVKDRUEER	RYTORFLAAY	BSEGSINTID	140
Ba	ch.	t	DWGDWTQKFT	I GORPSWONET	TELETVDWFE	HIDPLICAMIA	PYVKDKAEEN	BA BA	SADGQIRADO	137
OB.	Sh	1	PYWRDEILM	YFGALLYSEY	OLFNAHSSVO	ROCLEDTIRO	TAVFAALDEV	DHACMIQMER	LFIAKLVPGF	210
Ba	th	1	PTWRTSSCH	YWGAFLFNEY BB	GLPNARSQGA	REALSDVIRV	BLAPHOPDKI	DIAQMIQLER	GFLAKIVPGF	207
OH.	зb		DASTOVPER	WTTDFITSGA	RATVOEINOG	VODWIEILWA	GHAVMIATEG	QFARREFTOR	LATVYGDTLT	290
Ba	th	1	DESTAVPRAL	WINGEVYESA	ALAVEGLMOE	VEDWIEBAPS	VHAVYDALPO	QFVRREFFQR	LAPRFODELT	277
OB.	зъ		PFFTAQSQT	FOTTAGAIDD	LEVYCLANDS	EFGAENETFL	NAWTERYLAS	SVAALEDEVO	LYARVERSRA	350
Ba	th	1	PFFINGAQT	FOIARQUVOD	LYYNCLGODP	EFSDYNRTVN	BNWTGHWLEP BO	TIAALRDFMO	LFARLPAG	345
-	-	es.	ATDSAGVSEJ	I LORABODARI	DYA	DARBONIC BALL	a coverage			
Ba	th	ł	TTDKEEITA	LIEVVDDWIE	DYASAIDPKA	DRDQIVEAVI	AGLE			
γ	c	HA	IN							
OB	315		MARPEPTHON	I SIRTEWEART	ARLTSVDOAT	RFIQDERLAY	TSPFRASYDI	DVDYQYIERR	TEEKLSVLKT	70
Ba	th	4	MAR-LOINS	TRDAWVINI	ARVITLERAN	Y2 Y2	TIPFRISYEL	DEDYLWIEAR	Y3	69
OB	35	1	ERLPVADLI	T KATTGEDRAN	VEATWIAKIN	AARSKYEADO	INTEPROLYN	PPVLPVNVFL	RTDAALOTVL	140
Ba	th	1	RAFNEVDFR	H KTAFGEDARS	VLDOTVAIDO	AARDIOWEAEI	1HIGPROATS	PPINPVNYFL	DOERGLOTRL	139
OB	3b		MEIRPEDYY	TPLEOLARES	GVEVLHLQA					
Ва	th		MELFONLNYY	TPLEELSKOP	dvrvvh ∢ yøl>					

Fig. 3. Alignment of amino acid sequences of MMOH OB3b and MMOH Bath derived from the reported nucleotide sequences. Identical residues from both sequences are in black and the nonidentical residues are in red. Fe ligands are shown in cyan. Corrections to the amino acid sequence of MMOH OB3b determined by this work are shown in green above the aligned sequences. Secondary structural elements of MMOH OB3b are marked, with pairs of enclosing arrows, below the aligned sequences. All indicated elements are helices unless containing the letter "s" (β strands).

cellent $2F_o - F_c$ electron density covering part of this polypeptide segment. All the sequence variations observed can be explained through the insertion, deletion, or change of one or two base pairs.

Active site of MMOH

The dinuclear iron cluster is located between the helices αB , αC , αE , and αF (see Figs. 2 and 4). As can be seen in Figure 6A, the ligands for both irons have octahedral geometry. The irons are separated by 2.99 Å, consistent with the Fe-Fe distance observed in recent EXAFS studies on MMOH (Shu et al., 1996a), the -160 °C MMOH Bath structure (Rosenzweig et al., 1995), and model complex studies (Thich et al., 1976; Borer et al., 1982; Zang et al., 1994). The irons are bridged by two hydroxides (or one hydroxide and one water), as well as a carboxylato-bridge from Glu α 144. In the plane of the irons and the hydroxo ligands are four endogenous ligands: Glu α 114, His α 147, Glu α 209, and His α 246. Below this plane is the Glu α 144 bridge. Above the plane, Wat 3 and Glu α 243 complete the coordination sphere. As mentioned previously, Glu $\alpha 209$ is one of the residues where the observed sequence differed from that reported previously. Figure 2A and C shows the $2F_o - F_c$ electron density for the region around Glu $\alpha 209$.

The presence of a second bridging hydroxide (OH2) was indicated by a $5\sigma F_o - F_c$ feature. However, because the $2F_o - F_c$

Fig. 2. (*Facing page*) Selected sections of $2F_o - F_c$ electron density map of the final model of MMOH OB3b contoured at 1σ . **A:** Portion of the active site and the adjacent hydrophobic cavity occupied by four water molecules. **B:** Residues $\beta 363 - \beta 368$ of MMOH OB3b, a part of the 26 consecutive sequence corrections found in this region. **C:** A closeup view of the $2F_o - F_c$ (shown in light gray, contoured at 1σ) and $F_o - F_c$ electron density (shown in dark, contoured at 5σ) at $\alpha 209$ Glu. Side-chain atoms $C\gamma$, $C\delta$, $O \in 1$, $O \in 2$ for residue $\alpha 209$ were omitted from the refinement and from the map calculation.



Fig. 4. Structure of MMOH OB3b. A: Secondary structure of dimer with cylinders representing the helices and arrows representing the β strands. The crystallographic twofold axis is vertical. α chains are colored yellow and pink, β chains are colored gray and cyan, and γ chains are green. Iron atoms are shown as green spheres, and the hydroxides and the water molecules at the active site, as well as those at the adjacent hydrophobic cavity, are shown as red spheres. B: Stereo drawing of C_{α} trace of dimer colored as in A. C: Solvent-exposed surface colored by electrostatic potential (red, negative; blue, positive) as calculated by GRASP (Nicholls et al., 1991).

density for OH2 was less clear than that for OH1, OH2 was not added to the model until late in the refinement. At that point, $F_o - F_c$ density calculated after omitting the OH2 showed a peak >10 σ and the feature could be accounted for in no other way. It was also confirmed by the difference map calculated using the final model coordinates after omitting the two hydroxides by setting their occupancies to zero and a cycle of refinement that showed 11 σ peak at OH2 and 13 σ peak at OH1. OH2 is held less tightly than OH1 as evidenced by the longer distances to the irons (2.15 Å and 2.17 Å versus 1.71 Å and 2.04 Å), weaker $2F_o - F_c$ density, and a slightly higher temperature factor (14.8 Å² versus 11.5 Å²). It is not possible to distinguish between oxo, hydroxo, and aquo ligands from the electron density map. However, each of these potential bridging ligands have characteristic Fe-O bond lengths

Nucl. alt.:			0	3	3	3G	4	2		т	Δ									3	Δ.	с	GT										Δ		
Coden:	GA	GA	AG	rcg	cg	CGI	cod	BAC	ca	ATC	dC(GCC	GG	CG	FCT	00	GA	3 90	GC1	rge	A	↓ ¢CG	L TCT	TCO	agc	GAT	TGG	AG.	ATC	GAT	TA	CG	+	GA	CAAG
Rep. seq: Shiftl:	E R	K	s	R	RA	A .	R	T	R D	R	R	1	G	R V	L S	R	E	G A	A L	Å	2	R	S L	R	R	I L	GE	RD	SP	I	Т	R	P	D T	R
Shift2:	20	E	V	A		R,	G	P		1	A	P	A	. 1	g	P	R	P			5	۷	5		3 1	D	W 1	5	I	D	Y	A	R		2
Obs. seq:	E	R	V	7	G	A		\mathbf{T}	D	s	1	A.	C‡	V	ŝ		Ε	A	L	5	2	RV	5	6.3	1	D I	W 2	5	Ī.	D	Ý	А		D	ĸ

Fig. 5. Three reading frames of MMOH OB3b gene corresponding to residues β 346– β 375 are shown in lines "Rep. seq." "Shift1," and "Shift2." Sequence differences of the amino acids in the reported sequence (Cardy et al., 1991b) are shown in red in line "Rep. seq." Observed amino acids are shown in green. The corrected sequence from our study is also shown on the bottom line. Nucleotide sequence alterations required for the observed amino acid corrections are shown on top of the DNA strand. Δ stands for deletion of the nucleotide indicated. Nucleotide insertions are indicated by the other arrows.

that have been established from model studies. Bridging oxo ligands typically have Fe-O bond lengths of 1.7–1.9 Å (Kurtz, 1990), whereas the corresponding bond lengths for hydroxo and aquo bridging ligands are 1.9–2.1 Å (Armstrong & Lippard, 1984; Zang et al., 1994) and 2.15–2.3 Å (Hagen & Lachicotte, 1992), respectively. We assigned these two bridging atoms in the diiron cluster of MMOH OB3b as hydroxides based upon the observation that



Fig. 6. Geometry of the diiron center. A: Diiron center in MMOH OB3b. B: Superposition of diiron center of one of the monomers of MMOH Bath at 4°C (Rosenzweig et al., 1993) (shown in gray) onto that of MMOH OB3b at 18°C (shown in black).

their average bond lengths to the ferric ions are at the minimum (1.91 \AA) and maximum (2.15 \AA) distances, respectively, expected for this type of bridge. The longer bond might also be assigned to an aquo bridge, but the aquo bridge observed in the MMOH Bath diiron cluster in this position has a considerably longer average bond length (2.4 \AA) , suggesting that the bond in the current structure derives from a bridging hydroxo ligand. Moreover, the short Fe-Fe distance is more consistent with a hydroxo than an aquo bridge (Hagen & Lachicotte, 1992). The assignment of the shorter bonds as an oxo bridging ligand is disfavored on the basis that no optical spectrum is observed from the cluster in the spectral region expected for well-characterized oxo-bridged diiron clusters (Fox et al., 1989; Kurtz, 1990). Moreover, the exchange coupling observed for the cluster is much smaller than that of any known oxo-bridged diiron cluster (Fox et al., 1993).

Above and behind the diiron cluster, as oriented in Figure 6, is a hydrophobic cavity (volume 70 Å³) lined by Leu α 10, Gly α 13, Ala α 117, Ala α 120, Gln α 140, Cys α 151, Ala α 152, Ile α 171, Met α 184, Phe α 188, Phe α 192, Leu α 204, Thr α 213, Leu α 216, Ile $\alpha 217$, Phe $\alpha 236$, Val $\alpha 239$, Gln $\alpha 278$, and Phe $\alpha 282$. In this cavity, there is a region of $2F_o - F_c$ density that is not due to protein and has proven difficult to assign to a specific exogenous molecule (see Fig. 2A). During the refinement, we attempted to place several molecules in this density, including phosphate, because it was the only component in the crystallization solution of the appropriate size, but none were found to fit satisfactorily. Other obvious species that were placed initially in the density were methane and methanol; however, a single molecule was too small for the density and multiple molecules did not seem to fit the density very well. Because no species fit the density well, four solvent molecules (Wat 4, Wat 5, Wat 6, and Wat 7) were finally placed in the density. The average B-factor for these four water molecules is 40.0 Å², which is comparable to 37.1 Å², the average *B*-factor for all the water molecules in the structure.

Comparison with MMOH Bath

The high level of sequence identity of MMOH OB3b with MMOH Bath (see Fig. 3) suggests that similar structures will be observed, and this is the case. It should be mentioned here that the MMOH OB3b crystallizes as a monomer per asymmetric unit. Thus, the crystallographic twofold symmetry requires exact symmetry in the holoenzyme of MMOH OB3b. MMOH Bath crystallizes as a dimer in an asymmetric unit with noncrystallographic twofold symmetry restraints relating the monomers. It is not stated in either of the reports (Rosenzweig et al., 1993, 1995) or the PDB entry 1MMO whether noncrystallographic symmetry restraints were used during refinement of MMOH Bath. Superposition of the C_{α}s of the MMOH Bath monomers produces an RMS difference of 0.28 Å. Superposition of the two protomers of MMOH Bath found in its asymmetric unit with the crystallographic dimer found in the MMOH OB3b unit cell results in an RMS difference over all $C_{\alpha}s$ of 0.76 Å. Superposition of the individual monomers reduces this difference to 0.56 Å. The largest differences (those greater than 1.5 Å) between MMOH OB3b and MMOH Bath for the C_{α} trace are found at the terminal residues, and at residues $\alpha 340$, $\alpha 349$, $\alpha 430 - \alpha 435$, $\beta 32 - \beta 40$, $\beta 47$, $\beta 250 - \beta 280$, $\beta 325 - \beta 330$, and $\beta 338 - \beta 354$. The most significant changes observed are in the region β 345- β 354, corresponding to residues β 341- β 350 of MMOH Bath. This region is on the outer surface of the molecule and is not involved in any intermolecular contacts in either of the MMOHs. In MMOH Bath, the closest neighbor to this region consists of residues β 377– β 385 of the symmetry-related molecule, whereas in MMOH OB3b, the corresponding residues from the symmetry-related molecule are $\gamma 88 - \gamma 100$. The shortest distance between the molecule and the symmetry-related molecule at this interface for MMOH Bath is 3.93 Å, whereas that for MMOH OB3b is 3.68 Å. In both MMOHs, the C-terminal residue of the β chain is in this interface. In OB3b, there is a piece of continuous electron density that extends beyond the structurally derived last residue of the β chain (β 393) in the final model. However, the last two residues of the β chain (Lys β 394 and Asn β 395) could not be modeled due to very short contacts between the side chains of these two residues and the γ chain of the neighboring symmetry-related molecule. Hence, this region of density at the interface is modeled as four water molecules (Wat 699, Wat 729, Wat 730, and Wat 731).

The superposition of the structures of the diiron clusters of MMOH OB3b and that of one of the noncrystallographically related protomers of MMOH Bath (4 °C structure) are shown in Figure 6B. Overall, the active sites of both hydroxylases are quite similar. With the correction of Glu α 209, the protein ligand backbones agree with an RMS difference of 0.16 Å (averaged for both monomers of MMOH Bath). This difference is comparable to that seen between MMOH Bath protomers (0.10 Å). Superposition of the diiron cluster ligands and the residues that line the hydrophobic cavity gives an RMS difference for the backbone atoms between MMOH OB3b and MMOH Bath of 0.19 Å (averaged for both the monomers of MMOH Bath), which is again comparable to that between the monomers of MMOH Bath) (0.13 Å). With this level of structural similarity, the significantly higher specific activity for MMOH OB3b (Fox et al., 1989) cannot be explained easily by differences in active site structure. More specifically, the difference does not derive from the residue at position 209, as suggested previously by Rosenzweig et al. (1993), because this residue is shown here to be the same in both enzymes.

The diiron clusters of the two MMOHs do exhibit some significant differences in distance between the iron atoms and the exogenous bridging ligands. Table 2 lists the bond lengths and angles of the active site structure of MMOH OB3b for both form 1 and 2, and those of the two protomers of MMOH Bath at 4 °C and -160 °C (Rosenzweig et al., 1993, 1995). The major differences between the 4 °C MMOH Bath diiron cluster structure and that of MMOH OB3b is the longer Fe-Fe distance and the presence of an acetate (rather than a hydroxide) as the second exogenous bridge in the MMOH Bath cluster. In contrast, in the MMOH Bath structure determined at -160 °C (Rosenzweig et al., 1995), the Fe-Fe distance and exogenous bridging ligands are in excellent agreement with those seen for MMOH OB3b. Rosenzweig et al. (1995) argue that the geometry seen in the $-160\,^\circ C$ structure might be the result of including glycerol in the mother liquor as a cryoprotectant; but the results reported here for MMOH OB3b suggest that this is not the case. The similarity of the -160 °C MMOH Bath and the 18 °C MMOH OB3b structures suggests that the Fe₂-O₂ diamond core that they have in common may be a general feature of MMOHs.

Recent evidence suggests similarities in structure and reactivity between synthetic model compounds containing high valent Fe₂O₂ diamond core dinuclear iron complexes (Dong et al., 1995a, 1995b; Que & Dong, 1996) and the reactive high valent intermediate X of the R2 subunit of ribonucleotide reductase and compound Q of the MMOH catalytic cycle (Lee et al., 1993a, 1993b; Sturgeon et al., 1996). This resemblance suggests that the ability to form such a structure is important for the oxygen activation and insertion reactions that are catalyzed. On the other hand, EXAFS studies of the diferric MMOH OB3b (Shu et al., 1996a) indicate that, in frozen solution, about 40% of clusters have a 3.4-Å Fe-Fe distance, similar to the 4 °C MMOH Bath structure, whereas the remainder have a 3.0-Å Fe-Fe distance, similar to the -160 °C MMOH Bath and 18 °C MMOH OB3b structures. Consequently, all of the structures determined thus far for MMOH may be valid for the enzyme in solution under certain circumstances and may be indicative of the inherent flexibility of the cluster. Such flexibility may be nec-

	OF	33b		
	Form 1	Form 2	В	ath
Temperature	18 °C	18 °C	4°C	-160 °C
Fe-Fe distance	2.99 Å	3.07 Å	3.46 Å, 3.36 Å	3.04 Å, 3.14 Å
First bridge	OH	OH	OH	OH
Fe1-OH	1.71 Å	1.79 Å	1.71 Å, 1.69 Å	1.78 Å, 1.62 Å
Fe2-OH	2.04 Å	2.18 Å	1.98 Å, 2.06 Å	1.94 Å, 2.02 Å
Angle Fe1-OH-Fe2	105.4°	100.8°	138.6°, 127.7°	109.0°, 120.0°
Second bridge	ОН	ОН	CH3COOH	H ₂ O
Fe1-OH	2.15 Å	1.93 Å	2.56 Å, 1.95 Å	2.43 Å, 2.24 Å
Fe2-OH	2.17 Å	2.45 Å	2.30 Å, 2.05 Å	2.58 Å, 2.51 Å
Angle Fe1-OH-Fe2	87.6°	88.1°	N/A	Not reported

Table 2. Bond distances and angles of the diamond core found in MMO hydroxylases

^aBath data for the 4 °C structure are derived from the PDB entry 1MMO, and those for -160 °C are from Rosenzweig et al. (1995).

essary to accommodate various intermediates of the catalytic cycle that appear to have quite different diiron core structures. These intermediates range from the proposed peroxo-bridged compound P, which may require an Fe-Fe distance of up to 4 Å (Kim & Lippard, 1996), to the proposed diamond core bridged compound Q, in which the Fe-Fe distance is 2.47 Å (Shu et al., 1996b).

Electron density maps for both MMOH OB3b and MMOH Bath at -160 °C indicate the presence of some exogenous molecule(s) in the active site of MMOH. In the case of MMOH Bath, this density was interpreted as that of acetate, because acetate was present in the crystallization solutions (Rosenzweig et al., 1992). It is doubtful that acetate is the component present in the active site of MMOH OB3b because acetate is not in any of the solutions used for protein purification or crystallization. As mentioned previously, the unassigned density has been filled tentatively by four water molecules; however, this density cannot be unambiguously assigned at present for either enzyme.

Comparison with cytochrome $P-450_{CAM}$ and other diiron proteins

MMOH has been compared widely to cytochrome P-450, because it also activates molecular oxygen for incorporation into many substrates containing unactivated C-H bonds. In both enzymes, iron centers are reduced to allow the binding of molecular oxygen. The oxygen is then cleaved to produce water and a high valent reactive iron-oxo species that is capable of breaking a stable C-H bond (Ortiz de Montellano, 1986; McMurry & Groves, 1986; Lipscomb, 1994). In the case of cytochrome P-450, it has been proposed that breaking the O-O bond of oxygen requires protons to be shuttled into the active site to form water from one of the oxygen atoms as the high valent reactive species is formed. If the mechanism of MMO is fundamentally similar to that of cytochrome P-450, as we have proposed (Fig. 1, Fox et al., 1989, 1991), then a similar residue might be expected to be present in the active site of MMO to facilitate proton transfer. Because the only residues with side chains capable of forming hydrogen bonds in the active site of MMOH are Thr α 213 and Cys α 151 (see Figs. 2A and 7A), one of these residues could be involved in the proton transfer. In cytochrome P-450_{CAM} (Poulos et al., 1985), Thr 252 Oy is 5.7 Å from the heme iron. This distance compares with 6.3 Å and 6.8 Å for Thra 213 Oy from Fe1 and Fe2 of MMOH, respectively. Figure 7B shows the active site of cytochrome P-450_{CAM} oriented to emphasize this similar geometry. Mutation of Thr 252 to Ala in cytochrome P-450 uncouples the ability of this enzyme to hydroxylate substrate, with little or no product being formed (Imai et al., 1989; Martinis et al., 1989). Comparison of the amino acid sequences of MMOH with the diiron containing enzymes toluene 4-monooxygenase, phenol hydroxylase, and the stearoyl-ACP desaturases (Fox et al., 1994) reveals that the residue corresponding to $\alpha 213$ is always a threonine.

It has been suggested that a cysteinyl radical at Cys $\alpha 151$ might play a role in the hydroxylation chemistry at the active site of MMOH (Nordlund et al., 1992; Rosenzweig et al., 1993; Feig & Lippard, 1994). A basis for this suggestion can be seen by comparing the diiron centers of MMOH and of the R2 subunit of ribonucleotide reductase. As seen in Figure 7C, the residue corresponding to Cys $\alpha 151$ of MMOH is Tyr 122 of ribonucleotide reductase R2, which is the site of the stable tyrosyl radical gener-



Fig. 7. Active sites of (A) MMOH OB3b, (B) cytochrome P-450_{CAM}, and (C) ribonucleotide reductase R2 protein. Iron is shown in red and the hydroxide is shown in green as spheres. Residues corresponding to Cys α 151 of MMOH OB3b are orange, those corresponding to Thr α 213 are green, and those ligating the iron are yellow. The solvent-accessible surface near the iron(s) is shown with a transparent cyan surface.

ated by reaction of the dinuclear iron cluster of the protein. However, the amino acid sequence of toluene 4-monooxygenase has a glutamine in the homologous position, whereas the stearoyl-ACP desaturases have leucine (Fox et al., 1994). This suggests that Cys α 151 may not be critical to catalytic function.

Effects of component B

As stated earlier, component B was included in the crystallization solution that generated crystal form 2. Component B, by forming a tight complex with MMOH, has many significant effects on the activity of MMO that were summarized in the introduction (Fox et al., 1991; Wallar & Lipscomb, 1996). Both steady state and transient kinetic studies show that optimal rate enhancements are achieved when the bound complex stoichiometry is 2 MMOB:1 MMOH, that is, 1 MMOB per protomer or active site. Despite the fact that MMOB was present in a 2:1 stoichiometry during crystallization and a new crystal form was obtained, the electron density map showed no density for component B in the crystal structure. Moreover, SDS-PAGE of washed form 2 crystals indicated that component B was not present.

Growth of the crystal form 2 required the presence of component B. Duplicate crystallization trials under the same conditions except that component B was absent did not produce crystals. Previous studies have indicated (1) that interaction of MMOB and MMOH alters the MMOH structure and (2) that these alterations reverse slowly after dissociation of the complex (Fox et al., 1991; Froland et al., 1992; Liu et al., 1995a; Y. Liu, J. Nesheim, K.E. Paulsen, M.T. Stankovich, & J.D. Lipscomb, manuscript submitted). This hysteretic effect may cause a general change in the structure of the population of MMOH in the presence of MMOB, which is required to nucleate crystal form 2. In form 2 crystal structure, the dimer is rotated by 13.2° around the crystallographic a axis, with a 1.2° rotation of one monomer relative to the other from the crystallographic dyad with respect to form 1. Superposition of the monomers observed in both crystal forms produces an RMS difference of 0.60 Å over all atoms. Because this difference is comparable to that observed in independent refinements of the same protein (Ohlendorf, 1994), the structural changes imparted by component B binding are either too subtle to be seen at 2.4 Å resolution or have decayed back to native state.

An examination of the solvent-exposed surface of MMOH (Fig. 4C) reveals a negatively charged depression approximately 35 Å in diameter to the left of the twofold axis and directly above helices A, E, and F. Helices E and F participate in the active site by supplying three iron ligands ($\alpha 209$, $\alpha 243$, and $\alpha 246$) and by forming half of the active site pocket occupied by Wat 699, Wat 729, Wat 730, and Wat 731. Helix E is bent into three fragments ($\alpha 197-\alpha 204, \alpha 208-\alpha 211$, and $\alpha 213-\alpha 226$). By binding to this region, component B could alter the iron ligand geometry, as well as the volume of the active site, and thus modulate activity.

Conclusions

The observation of a diamond core structure for the essential diiron cluster of the active site of MMOH OB3b similar to that reported for the -160 °C structure of MMOH Bath shows that this is the dominant structure that has been maintained across the family of sMMO enzymes from highly divergent methanotrophs. Moreover, the presence of the diamond core structure in the highly active *M. trichosporium* MMOH that has been crystallized and studied at near room temperature strongly supports this conclusion. The diamond core structure is probably lost during some stages of the catalytic cycle, but it appears to be the most likely structure for the critical transient intermediate, compound Q, that appears to insert oxygen into the unactivated C-H bond of methane. The significance of the compact diamond core structure to oxygen activation and insertion chemistry is currently under intense investigation

(Que & Dong, 1996). This novel structure presents many mechanistic possibilities beyond those considered in the past for heme oxygenases (Feig & Lippard, 1994; Wallar & Lipscomb, 1996). The structural insight provided by the current study supports the occurrence of such a species in MMOH catalysis and supplies critical metric information for ongoing biomimetic studies.

Materials and methods

Crystal form 1

The growth of M. trichosporium OB3b and the purification of MMOH were as reported previously (Fox et al., 1989, 1990a). The specific activity at 23 °C and iron content of MMOH OB3b are 800-1000 nmol/min/mg and 1.8-2.0 diiron clusters/mol, respectively. Diffraction-quality crystals were obtained as reported by a hanging drop method with repeated macroscopic seeding at 18 °C (Froland et al., 1994). Typical crystal size is 0.7 mm \times 0.7 mm \times 0.3 mm. The crystals are in space group C222₁, with one monomer per asymmetric unit. X-ray data were collected using a MAR imaging plate mounted on beamline X12C at the Brookhaven National Laboratory. Images were processed using DENZO (Otwinowski, 1993). Although diffraction to about 2.5 Å resolution was resistant to radiation damage, the higher-resolution data (to <1.8 Å resolution) had a lifetime of about 1 h on X12C. Consequently, a high-resolution native data set was formed using data collected from nine crystals of MMOH OB3b.

Coordinates of one protomer of MMOH Bath (Rosenzweig et al., 1993) (chains B, D, and G) were extracted from the PDB data file 1MMO. Using these coordinates, a cross rotation function was calculated with XPLOR (Brünger, 1992). Two searches, each with vector lengths of 45 Å and 35 Å, were conducted using data between 8 and 3.5 Å and 8 and 4 Å. Both searches yielded the same prominent peak that, following Patterson correlation refinement, gave a peak that was 9.7 times higher than the next highest peak. Coordinates corresponding to this peak were used in the translation-search by XPLOR. This resulted in a single solution and provided an initial model for MMOH OB3b. After setting aside 2% (1,721 reflections) of the observed data for the calculation of R_{free} , molecular dynamics refinement by simulated annealing using XPLOR reduced the R-factor and R_{free} for data between 8–2.8 Å, with $F > 2\sigma$, to 24.8% and 34.5%, respectively. Because the values of R_{free} and R indicated a correct solution, all the data were used in the subsequent refinements and map calculations. Group temperature factor refinement using the data between 8 and 3.5 Å resolution and $F > 3\sigma$ produced an *R*-factor of 21.7%. Using TOM (Jones, 1978), this model was examined and the sequence of the model was altered to that expected for MMOH OB3b (Cardy et al., 1991b). Several cycles of Powell minimization were run at varying resolutions trying to reconcile the model with the expected amino acid sequence. When this failed, residues that disagreed with the published sequence were replaced with either alanines or glycines and main-chain breaks were made at the points of residue insertions. Additional cycles of refinement accompanied by phase extension to 2 Å resolution with $F > 2\sigma$ converged to an R-factor of 22.2%. By this time, several of the randomly distributed sequence changes reported in Figure 3 had been made and clear density for solvent molecules was apparent. After adding 230 water molecules with a polyalanine model for β 345– β 372, the *R*-factor dropped to 19.0% for data between 5 and 2 Å, with $F > 2\sigma$. At this point, an examination of the reading frames for the β chain of MMOH produced an amino acid sequence that agreed with that seen in the electron density maps. In addition, the density now unambiguously indicated the presence of the second bridging hydroxide. Further refinement after these corrections and adding another 200 water molecules reduced the R-factor to 16.2% for data between 5 and 2 Å, with $F > 2\sigma$. During the next cycles of map examination, addition of solvent molecules, and refinement, various models (a phosphate, two methanols with half occupancies each occupying alternating confirmations, two methanes, an ethanol, an ethane, a serine, a cysteine) were tested in the density feature found in the active site. Because no single model explained this density adequately, a group of four solvents was placed in the model, which, on refinement, produced an R-factor of 14.4% for data between 5 and 2 Å, with $F > 2\sigma$. Additionally, two water molecules were found on special positions. On inclusion of these water molecules, refinement produced an R-factor of 13.7% for the final model for data between 5 and 2.0 Å, with $F > 2\sigma$ (14.5%) with no cut off).

Crystal form 2

The growth of *M. trichosporium* OB3b and purification of MMOH and MMOB were as reported previously (Fox et al., 1989, 1990a). The specific activity at 23 °C of MMOB was 7,000 nmol/min/mg. Single form 2 crystals were obtained by vapor diffusion starting with a 10 mg/mL mixture of MMOH OB3b and recombinant MMOB with a molar ratio of 2 MMOB:1 MMOH OB3b. The reservoir solution consisted of 18% (w/v) PEG₃₃₅₀, 167 mM Na₂HPO₄, and 10 mM citric acid at pH 7.5. After combining 7 μ L of the MMOH/component B mixture with 7 μ L of the mother liquor (4% PEG₃₃₅₀ and 100 mM phosphate/citrate buffer, pH 7.7), large single crystals were obtained after 16 days. The crystals are in space group C222₁, with one monomer per asymmetric unit. A native data set was collected to 2.71 Å resolution using a Siemens/ Xentronics X100 area detector mounted on a Rigaku rotating anode source.

A nearly completely refined model of MMOH OB3b was used as a search model for molecular replacement in the program XPLOR (Brünger, 1992). The rotational search produced a peak that, after Patterson correlation refinement, was 8.1 times higher than the next highest peak. Coordinates corresponding to this peak were used in the translation search by XPLOR. This resulted in a single solution and provided an initial model for the form 2 of MMOH OB3b. After setting aside 5% (1,658 reflections) of the observed data for the calculation of R_{free} , molecular dynamics refinement by simulated annealing using XPLOR produced a model with an *R*-factor of 19.2% and R_{free} of 29.3% for data between 8 and 2.8 Å, with $F > 2\sigma$. Because the values of R_{free} and R indicated that the solution was correct, all the data were used for further refinement and map calculations. Group temperature-factor refinement using data between 8 and 2.7 Å and $F > 2\sigma$ brought down the *R*-factor to 17.3%. Powell minimization using XPLOR produced a model with an R-factor of 16.7% for data between 8 and 2.7 Å, with $F > 2\sigma$. Examination of the electron density map, addition of the two Fe³⁺, the two bridging hydroxides, and the exogenous ligand (one water molecule) to the iron, and Powell minimization using XPLOR produced a model with a crystallographic R-factor of 16.3% using 22,264 unique reflections between 5 and 2.7 Å resolution, with $F > 2\sigma$. B-factors were found as low as 2 Å² for many of the side-chain atoms. Hence, B-factors for the whole model were reset to the Wilson B-factor before the next cycle of refinement. At this stage, the two bridging hydroxides had poor $2F_o - F_c$ density, which we attribute to the lower-resolution data of the diffraction. Even though the $2F_o - F_c$ as well as positive $F_o - F_c$ density was obvious for about 240 water molecules, none were placed in the model except one water molecule at the hydrophobic cavity because of the limited resolution. The complete model was checked and a final set of positional and temperature-factor refinement produced an *R*-factor of 15.2% for the final model for data between 5 and 2.7 Å, with $F > 2\sigma$ (17.8% with no cut off).

During refinement of both crystal forms of MMOH OB3b, NOE square well constraints were placed on the irons and their ligands. For crystal form 1, the Fe-ligand NOEs were set to 2.0 ± 0.4 Å and the Fe-Fe NOE was set to 3.0 ± 0.4 Å. For crystal form 2, the NOEs were set to the values observed in crystal form 1 (± 0.4 Å). Coordinates of both forms of the crystal structure of MMOH OB3b have been submitted to Protein Data Bank with accession numbers 1MHY and 1MHZ.

Figures 2, 4A, 4B, and 6 were made using SETOR (Evans, 1993). Figures 4C and 7 were made using GRASP (Nicholls et al., 1991).

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