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Coupling oxygen consumption with hydrocarbon oxidation in bacterial multicomponent monooxygenases

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Abstract

A fundamental goal in catalysis is the coupling of multiple reactions to yield a desired product. Enzymes have evolved elegant approaches to address this grand challenge. A salient example is the biological conversion of methane to methanol catalyzed by soluble methane monooxygenase (sMMO), a member of the bacterial multicomponent monooxygenase (BMM) superfamily.

sMMO is a dynamic protein complex of three components: a hydroxylase, a reductase, and a regulatory protein. The active site, a carboxylate-rich non-heme diiron center, is buried inside the 251-kDa hydroxylase component. The enzyme processes four substrates: O₂, protons, electrons, and methane. To couple O₂ activation to methane oxidation, timely control of substrate access to the active site is critical. Recent studies of sMMO, as well as its homologs in the BMM superfamily, have begun to unravel the mechanism. The emerging and unifying picture reveals that each substrate gains access to the active site along a specific pathway through the hydroxylase. Electrons and protons are delivered via a three-amino acid pore located adjacent to the diiron center; O₂ migrates via a series of hydrophobic cavities; and hydrocarbon substrates reach the active site through a channel or linked set of cavities. The gating of these pathways mediates entry of each substrate to the diiron active site in a timed sequence and is coordinated by dynamic interactions with the other component proteins. The result is coupling of dioxygen consumption with hydrocarbon oxidation, avoiding unproductive oxidation of the reductant rather than the desired hydrocarbon.

To initiate catalysis, the reductase delivers two electrons to the diiron(III) center by binding over the pore of the hydroxylase. The regulatory component then displaces the reductase, docking onto the same surface of the hydroxylase. Formation of the hydroxylase-regulatory component complex (i) induces conformational changes of pore residues that may bring protons to the active site; (ii) connects hydrophobic cavities in the hydroxylase leading from the exterior to the diiron active site, providing a pathway for O₂ and methane, in the case of sMMO, to the reduced diiron center for O₂ activation and substrate hydroxylation; (iii) closes the pore, as well as a channel in the case of four-component BMM enzymes, restricting proton access to the diiron center during formation of “Fe₂O₂” intermediates required for hydrocarbon oxidation; and (iv) inhibits undesired electron transfer to the Fe₂O₂ intermediates by blocking reductase binding during O₂ activation. This mechanism is quite different from that adopted by cytochromes P450, a large class of heme-
containing monooxygenases that catalyze very similar reactions as the BMM enzymes. Understanding the timed enzyme control of substrate access has implications for designing artificial catalysts. To achieve multiple turnovers and tight coupling, synthetic models must also control substrate access, a major challenge considering that nature requires large, multimeric, dynamic protein complexes to accomplish this feat.

**Graphical Abstract**

![Graphical Abstract](image_url)

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**1. Introduction**

How can reactions among multiple substrates be coupled to generate a desired product? This challenge is frequently seen in biocatalysis, especially in achieving the most difficult chemical transformations. One example is the biological activation of inert C–H bonds. This transformation is catalyzed by several metalloenzymes, including the heme-containing cytochromes P450,[1–4] the dicopper-containing particulate methane monooxygenase,[5,6] and the family of non-heme diiron-containing bacterial multicomponent monooxygenases (BMMs).[7–10] These enzymes couple reactions involving four substrates (eq 1), namely, oxygen, protons, electrons, and a hydrocarbon RH.

\[
O_2 + 2H^+ + 2e^- + RH \rightarrow H_2O + ROH \quad (1)
\]

Timed control of substrate addition to the active site is essential for coupling \( O_2 \) activation to substrate hydroxylation. Without proper control, the following uncoupled reactions can occur (eq 2, 3):[2,11,12]

\[
O_2 + 4H^+ + 4e^- + RH \rightarrow 2H_2O + RH \quad (2)
\]

\[
O_2 + 2H^+ + 2e^- + RH \rightarrow H_2O_2 + RH \quad (3)
\]

Such uncoupling could be detrimental to the organism, for example, by wasting reducing equivalents and by generating hydrogen peroxide, a reactive oxygen species.

Recently, we have come to understand that the desired coupling (eq 1) involves carefully programmed access of substrates to the diiron active sites in the hydroxylase components of soluble methane monooxygenase (sMMO) and its homologs in the bacterial multicomponent monooxygenase (BMM) superfamily. These enzymes are of great interest in the context of the global carbon cycle[13] and bioremediation.[14,15] They are complex systems typically comprising three to four protein components.[8,16] The flagship enzyme, sMMO, has three
components, a hydroxylase (MMOH), a reductase (MMOR), and a regulatory protein (MMOB). The 251-kDa hydroxylase is a homodimer with three subunits (α, β, and γ) in each protomer. Each α subunit hosts a diiron center, where dioxygen is activated and substrate hydroxylated. The 38-kDa reductase is responsible for providing two electrons during each round of the catalytic cycle. It transfers electrons from the reduced form of nicotinamide adenine dinucleotide (NADH) to the diiron center, via its flavin adenine dinucleotide (FAD) cofactor in the C-terminal domain and an [Fe₂S₂] cluster in the ferredoxin domain. The third component is a 16-kDa regulatory protein, which accelerates dioxygen activation and methane oxidation and is critical for coupling. Some members of the BMM superfamily, such as toluene/o-xylene monooxygenase (ToMO) and toluene-4-monooxygenase (T4mo), require an additional Rieske-type [Fe₂S₂] protein for reducing the hydroxylase. Dynamic interactions among these component proteins control substrate access to achieve some of the most challenging chemical conversions.

Since our previous Account published in 2007, we have learned that coupling in BMM enzymes relies critically upon the control of substrate access to the catalytic diiron center. Here we describe the entry pathway for each substrate as well as the molecular details of the mechanism used to control substrate access. Finally, we compare the tactics used by BMM enzymes with those adopted by P450s, a large group of related monooxygenases.

2. Control of electron and proton transfer: gating the pore

Accurate control of electron and proton transfer is vital for tight coupling. Electron transfer initiates catalysis by reducing the diiron(III) center in the resting hydroxylase to diiron(II) (Scheme 1). In subsequent steps, however, electron transfer is undesired, because extra electrons could quench the activated oxygen species in P*, Hperoxo, and Q. Proton transfer is required for O₂ activation, promoting heterolytic cleavage of the O–O bond in sMMO catalysis to generate the methane-oxidizing intermediate Q. Excess proton transfer could quench the activated oxygen species, however, leading to production of H₂O and H₂O₂ (eqs 2, 3). Accumulating evidence indicates that electrons and protons are transmitted through the pore (Figure 1a,b) and are gated through the interaction with the regulatory component.

2.1. The electron transfer pathway

The reduction of the diiron(III) center is mediated by the reductase component(s). To understand the mechanism of electron transfer control, we first need to identify the reductase binding site on the hydroxylase and the electron transfer pathway. As the shortest entry from the protein exterior to the diiron center, the pore is the leading candidate for the reductase binding site.

Recent studies of sMMO, T4mo, and ToMO all confirm this proposal. In the case of sMMO, the reductase binding site on the hydroxylase was determined by hydrogen-deuterium exchange mass spectrometry (HDX-MS). The data indicated that the ferredoxin domain of the reductase binds to a shallow region at the dimer interface of the hydroxylase, known as the “canyon”, covering the pore (Figure 1a). A computational docking study supported this conclusion (Figure 1c) and revealed that the gating residue of the pore, Glu240, lies...
nearly on a line between the \([\text{Fe}_2\text{S}_2]\) cluster of the MMOR ferredoxin domain and the diiron center of MMOH (Figure 1d).\(^{35}\) The importance of the pore residues Asn202 and Gln228 in ToMO (Asn214 and Glu240 in sMMO) for reductase binding was demonstrated in a biochemical study of ToMO.\(^{32}\)

Atomic details of the hydroxylase-reductase complex were revealed in an X-ray crystallographic investigation of T4MO.\(^{36}\) Consistent with our findings with sMMO\(^{35}\) and ToMO,\(^{32,37}\) the reductase (T4moC) covers the pore region in the complex (Figure 1e). The crystal structure further revealed a 12-Å distance between the \([\text{Fe}_2\text{S}_2]\) cluster of the reductase and the diiron center of the hydroxylase, favorable for biological electron transfer.\(^{38}\) The two metal clusters are linked through a hydrogen-bonding network, which connects the \([\text{Fe}_2\text{S}_2]\) cluster ligand His68 with the diiron center ligand Glu231, mediated by the pore residue Gln228 (corresponding to Glu240 in MMOH) (Figure 1f).\(^{36}\)

### 2.2. Control of electron transfer: binding competition at the pore

An understanding of the mechanism by which the BMM proteins control the timing of electron transfer required information about interactions between the hydroxylase, reductase, and regulatory components. Two models were considered, a non-competitive model whereby the regulatory component and the reductase bind to different sites on the hydroxylase\(^{12}\) and a competitive model where they compete for the same binding site on the hydroxylase. Recent structural studies strongly favor the latter. X-ray crystallographic results for the hydroxylase-regulatory protein complex revealed that the regulatory component binds to the canyon, the same region where the reductase binds, covering the pore. This conserved binding mode occurs in sMMO (Figure 2a),\(^{31}\) T4mo (Figure 2b),\(^{29}\) and PH (Figure 2c).\(^{39}\)

Multiple biochemical/biophysical characterization experiments further support the binding competition. In the case of sMMO, MMOB inhibits MMOR cross-linking to MMOH in a dose-dependent manner. The MMOR ferredoxin domain protein can displace MMOB from MMOH, as observed in a titration experiment by monitoring fluorescence anisotropy of MMOB carrying a fluorescent label.\(^{35}\) Binding competition between the regulatory component and the reductase was also reported for T4mo.\(^{36}\) Consistent with the competitive model, the regulatory component retards electron transfer from the reductase to the diiron center, an effect observed in sMMO\(^{20,35,40}\) and ToMO.\(^{37}\)

Binding competition between the reductase and the regulatory components at the pore region of the hydroxylase provides a basis for electron transfer control. By fine-tuning the binding affinity, the regulatory component could inhibit reductase binding during \(\text{O}_2\) activation, blocking undesired electron transfer to the diiron center. Indeed, the regulatory component exhibited increased binding affinity to the hydroxylase after reduction of the diiron center in sMMO isolated from *Methylococcus capsulatus* (Bath).\(^{41}\)

### 2.3. Proton and water transfer through the pore

Oxygen activation requires protons.\(^{10,26,27}\) As the only hydrophilic entry to the diiron center, the pore provides the route for proton transfer. Biochemical study of ToMO indicated
pore residue Thr201 to be critical for proton transfer during dioxygen activation.\textsuperscript{34} Kinetic isotope effects and pH profiles suggested that another pore residue, Gln228, mediates proton ingress to, and water egress from, the active site.\textsuperscript{32}

Structural studies revealed the molecular mechanism of proton transfer. In the case of sMMO, crystal structures showed Glu240 to be the gating residue in the pore, playing a key role in proton transfer.\textsuperscript{31} In the absence of other component proteins, this residue is hydrogen bonded to a water or hydronium ion on the surface of the hydroxylase.\textsuperscript{31,42} In response to the binding of the regulatory component, the carboxylic side chain of Glu240 moves inward\textsuperscript{31} in a manner suggesting a role in delivering a proton to the active site for O\textsubscript{2} activation and, in the process, closes down the pore (Figure 3a) to block undesired water/hydronium ion ingress that would quench reactive intermediates (Scheme 1).\textsuperscript{31} A similar conformational change may occur when the reductase binds to the hydroxylase. Thus Glu240 provides the basis for proton-coupled electron transfer.\textsuperscript{35}

Structures of protein complexes of T4mo revealed additional details. As observed in sMMO, the regulatory component and the reductase can both induce inward movement of the pore gating residue Gln228.\textsuperscript{29,36} A water molecule not observed in the free hydroxylase appeared in the pore in the hydroxylase-regulatory component (Figure 3b) and hydroxylase-reductase (Figure 1f) complexes, providing a putative mechanism for proton transfer. In the case of the reduced hydroxylase-regulatory component complex, this water molecule (HOH5) is close to the open coordination site for O\textsubscript{2} binding (Figure 3b), serving as an likely proton source for dioxygen activation.\textsuperscript{29} The crystal structures also revealed an important role of the conserved Thr201 in stabilizing the hydrogen bonding network at the active site,\textsuperscript{26,34} providing a structural basis for the importance of this residue in proton transfer and dioxygen activation.

3. Control of O\textsubscript{2} access

After reduction of the hydroxylase, O\textsubscript{2} needs to be activated by the reduced diiron center for substrate hydroxylation (Scheme 1). How does O\textsubscript{2} reach the buried diiron center, and how is the access controlled?

3.1. Cavities as the O\textsubscript{2} ingress route

Hydrophobic gas molecules, including O\textsubscript{2} and CH\textsubscript{4}, find their path to the diiron center through a series of cavities that extend over 36-40 Å through the protein interior. Multiple lines of evidence support this pathway. Structural characterization of Xe-pressurized sMMO and PH hydroxylases revealed xenon atoms, which are similar to O\textsubscript{2} molecules in terms of size, hydrophobicity, and polarizability,\textsuperscript{43} localized in these cavities. Halogenated alkanes, substrate analogs for sMMO bearing heavy atoms, can also occupy these cavities.\textsuperscript{44} Mutagenesis of residues in each of the three hydrophobic cavities in ToMO altered the dioxygen diffusion rate in a manner that correlated with the size of the cavities.\textsuperscript{45} These cavities are observed in all crystallographically characterized BMM hydroxylases and are likely to be conserved across the superfamily.\textsuperscript{30}
3.2. Control of O$_2$ access by the regulatory component

An interesting feature of these cavities was evident from the early studies. In several cases, cavities 1 and 2 are disconnected (Figure 4a), raising the question of how gas molecules could migrate through such a disconnected pathway. The crystal structure of the hydroxylase-regulatory component complex of sMMO provided critical insight. It revealed that, upon binding of the regulatory component MMOB, cavities 1 and 2 become connected (Figure 4b). MMOB residues Ser111 and Tyr8, at the interface with MMOH in the complex, trigger allosteric changes that alter the conformation of Phe188, the gating residue between the two cavities (Fig. 4).

The crystal structure also explains the importance of the N-terminal tail of MMOB. The regulatory components of ToMO, T4mo, PH, and most other members of the BMM superfamily do not have such tail, but in the case of sMMO, enzyme activity is severely compromised when the N-terminal tail is truncated or even partially truncated. The N-terminal tail is disordered in free MMOB, as revealed by solution NMR spectral studies, but it adopts an unusual ring-like conformation when bound to the surface of MMOH (Fig. 2a). A key residue in this tail in sMMO is Tyr8, which triggers allosteric changes in MMOH that lead to the changes depicted in Figure 4, with assistance from residue Ser111, which resides in the core of the regulatory protein.

Another important finding is that the function of MMOB in facilitating O$_2$ ingress depends on the oxidation state of the diiron center. In a study of the MMOH-MMOB complex in solution, the conformation of the MMOB N-terminal tail was assessed using nitroxide spin-labeled MMOB mutants by double electron-electron resonance spectroscopy. The data revealed that the N-terminal tail switches from a flexible to an ordered conformation in response to reduction of the diiron center from the diiron(III) to the diiron(II) state. The allosteric alterations observed crystallographically in the MMOH-MMOB complex are therefore not occurring for the diiron(III) form of the hydroxylase in solution, because Tyr8, a critical residue that induces allosteric changes in MMOH, is disordered and lacking a stable contact with the hydroxylase. Thus oxygen, and most likely the structurally similar hydrocarbon substrate methane, do not have access to the active site until the diiron center is reduced to diiron(II), when the N-terminal tail becomes ordered and Tyr8 forms a stable interaction with MMOH. Through such a mechanism, access of O$_2$ and CH$_4$ to the active site is coupled to reduction of the diiron center.

4. Hydrocarbon substrate ingress and product egress

The small hydrocarbon CH$_4$ finds its route to the active site through cavities in sMMO. Some BMM enzymes, however, process much larger, and in some case polar, hydrocarbons. In addition, the hydroxylation products are all more hydrophilic than the original substrate. What are the pathways for trafficking these substrates and products to and from the active site?

4.1. ToMO/T4MO: through the channel and the cavities

A prominent feature of the four-component toluene monooxygenases ToMO and T4mo is a 6 – 10 Å-wide, 35 Å-long channel in their hydroxylase components (Figures 5a, b).
extends from the protein surface to the active site and merges with cavities 1 and 2 at their junction.\textsuperscript{28,29} The channel was proposed as the substrate and product migration pathway in toluene monooxygenases (TMOs), because the product analog bromophenol occupied the channel in a crystal structure of ToMO.\textsuperscript{28} The possibility that this channel might also serve as the O\textsubscript{2} access pathway was unsupported by a mutagenesis study of ToMO, where variants with narrowed or widened channels did not affect the rate of O\textsubscript{2} diffusion to the diiron center.\textsuperscript{45} The cavities, which exist in TMOs as the O\textsubscript{2} migration pathway, can also provide aromatic substrates access to the active site.\textsuperscript{50,51} Additional studies indicated the cavities to be preferred as the substrate ingress pathway, whereas the more hydrophilic channel was the preferred product egress pathway.\textsuperscript{51}

The opening of the channel is also controlled by the regulatory component. Upon formation of the hydroxylase-regulatory component complex in T4mo, the channel collapses, making it inaccessible to small molecules (Figure 5c).\textsuperscript{29} This effect blocks solvent access to the active site during dioxygen activation, thereby increasing the coupling efficiency. Binding of the reductase to the hydroxylase leaves the channel open.\textsuperscript{36}

4.2. PH: through the pore or the cavities?

Similar channels occur in the hydroxylase components of PH and sMMO as revealed by computational analysis, but they are more constricted and less likely to be functional.\textsuperscript{30} In the case of PH, the migration pathways for the phenolic substrate and product are unclear. One possibility is through the cavities, because halogenated alcohols of different sizes could be located by X-ray diffraction in the cavities of the sMMO hydroxylase co-crystals,\textsuperscript{52} but direct evidence for such is lacking in the case of PH. The pore is an alternative option, considering its hydrophilicity and the larger opening (6 Å diameter) than the pores that occur in other BMM enzymes.\textsuperscript{39}

5. Similar function, different strategies: a comparison with cytochromes P450

The heme-containing cytochrome P450 monooxygenases catalyze the same overall reaction as BMM enzymes (eq 1), but are distinct in many respects. P450s are 40-55 kDa single-polypeptide proteins. Most of them do not require a regulatory component, also referred to as an effector protein, except for a few cases including P450cam, the reductase of which also serves as the effector.\textsuperscript{53} Moreover, P450s adopt very different tactical approach to controlling substrate access to the catalytic heme active site.

In the electron transfer step, delivery of the first electron to the heme iron center is gated by substrate binding, which prevents wasteful discharge of the reductant in the absence of substrate.\textsuperscript{1,2,4} Such a gating effect is achieved by displacing a water ligand from the heme iron upon substrate binding; as a result, the heme iron switches from low-spin to high-spin with a concomitant increase in the redox potential, which favors reduction.\textsuperscript{54-57} This control mechanism is not observed in BMM enzymes. In the subsequent O\textsubscript{2} activation step for P450s, an Fe(IV)=O species known as Compound I\textsuperscript{2,58} forms by oxygenation of the reduced heme site and subsequent e\textsuperscript{-}/2H\textsuperscript{+} transfer. Quenching of Compound I (eq 2) by further
electron transfer is disfavored over substrate oxidation as a result of the close proximity of substrate to this oxidizing species.\textsuperscript{2,59}

Control of proton and water access to the active site is also important for P450s. Without proper control, the peroxy intermediate that precedes formation of Compound I is quenched,\textsuperscript{2} releasing H\textsubscript{2}O\textsubscript{2} (eq 3). The hydrophobicity of the active site\textsuperscript{2,59} as well as the conserved alcohol-acid pair, residues Thr252 and Asp251 in P450cam,\textsuperscript{2,4,60} are essential for proper proton transfer. In addition, substrate binding expels undesired water molecules from the active site\textsuperscript{61} and, in many cases, induces a closed conformation that shields the active site from exposure to exterior solvent. This conformational change has been observed in P450cam\textsuperscript{61-63} and in P450s processing specific substrates,\textsuperscript{64-67} but may not be relevant to P450s processing more diverse substrates.\textsuperscript{68} Such global conformational dynamics are also critical for substrate specificity.\textsuperscript{62} In contrast, BMM enzymes use the regulatory component to block undesired proton and water access during O\textsubscript{2} activation, by closing the pore as well as the channel in the case of TMO. In addition, the hydroxylases of BMM enzymes do not undergo any global conformational changes.

6. Conclusions

BMM enzyme systems form dynamic complexes among their protein components to couple reactions of multiple substrates by timely control of their access to the active site. To achieve such control, different reactants are routed through different pathways in the hydroxylase. Electrons and protons traverse the pore; O\textsubscript{2} diffuses through the cavities; and hydrocarbon substrates use either the cavities or a special channel. The regulatory component gates access to these pathways by reversible binding to the canyon region, which includes the pore, in a manner that can be regulated by the oxidation state of the diiron center. In a catalytic cycle, the reductase transmits two electrons to the diiron(III) center by docking into the canyon. Next, the regulatory component displaces the reductase and inhibits excess electron transfer during O\textsubscript{2} activation by physically occluding the reductase. Through binding to the sMMO hydroxylase, the regulatory component restricts water and proton access by closing the pore, and, in the case of T4mo, it collapses a channel accessed by the aromatic substrates. Within all BMM hydroxylases, allostERIC changes induced by binding of the regulatory proteins connect the cavities, facilitating O\textsubscript{2} (and CH\textsubscript{4} in the case of sMMO) access to the diiron center. At the conclusion of a full catalytic cycle, the enzyme returns to the diiron(III) state, which lowers the binding affinity of the regulatory component to the diiron center. The reductase is then able to compete with the regulatory component for binding to the canyon, reduce the diiron center, and initiate the next round of catalysis.

Although a unified mechanism is emerging, there are some important outstanding issues. For instance, to fully support the proposed mechanism of electron transfer control, it would be valuable to have direct evidence that activated oxygen species can be quenched in the absence of the regulatory component or, by excess reductase that could displace the regulatory component. In addition, each BMM family member has uniquely characteristic properties. An example is the unique ability of sMMO to oxidize methane, which may reflect the apparent inability to generate intermediate Q in any other enzyme. This property
may derive from an as yet unknown distinctive strategy involving control of substrate access to the diiron center.

The present understanding of these control mechanisms has the potential to guide the engineering of improved biocatalysts for hydrocarbon oxidation using atmospheric oxygen. It also poses a great challenge for synthetic modeling of enzymes – how do we achieve substrate access control in synthetic models, something that nature accomplishes with the invention of large and complex proteins and protein complexes?

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Biography

Stephen J. Lippard is the Arthur Amos Noyes Professor of Chemistry at the Massachusetts Institute of Technology. His research areas lie at the interface of inorganic chemistry and biology.

Weixue Wang received his PhD in Biophysics and Computational Biology from the University of Illinois at Urbana-Champaign under the direction of Eric Oldfield. He was a postdoctoral associate with Stephen J. Lippard at the Massachusetts Institute of Technology. This year he assumed a position at the Janssen Pharmaceutical Companies of Johnson & Johnson.

Alexandria Deliz Liang received her Bachelor of Arts in chemistry from New College of Florida in 2010, where she worked in the laboratory of Suzanne E. Sherman. In 2010, Alexandria matriculated to the Massachusetts Institute of technology, where she recently completed her PhD, investigating the reactivity of toluene/o-xylene monooxygenase in the laboratory of Professor Lippard.

Reference


Scheme 1.
The catalytic cycle of sMMO. $R_{\text{red}}$ and $R_{\text{ox}}$ represent the reduced and oxidized reductase MMOR, respectively, and B the regulatory component MMOB.
Figure 1.
The reductase binding site on the hydroxylase. a) Structure of sMMO hydroxylase (PDB ID 1MTY), showing an area that is within 12 Å of the diiron center in one of the monomers (red) and the reductase binding site suggested by HDX-MS (orange). b) A close-up view of the pore in sMMO hydroxylase, showing the three residues that define the pore and Fe1 of the diiron center. c) Computationally docked model of the hydroxylase-reductase ferredoxin domain protein complex of sMMO. d) A close-up view of the binding interface in the docked model shown in c), viewed from the top, showing the [Fe₂S₂] cluster, the diiron center, and selected residues of the hydroxylase. e) Crystal structure of the hydroxylase-reductase complex of T4mo (PDB ID 4P1B). f) A close-up view of the binding interface in the docked model shown in e), viewed from the top.
Figure 2.
Crystal structures of the hydroxylase-regulatory component complexes of a) sMMO (PDB ID 4GAM), b) T4mo (PDB ID 3DHI), and c) PH (PDB ID 2INP). The hydroxylases are shown in cyan, and regulatory components in magenta.
Figure 3.
Regulatory component induced conformational changes at the pore. a) A close-up view of the pore in the hydroxylase-regulatory component complex of sMMO (PDB ID 4GAM), where the pore is closed. b) A close-up view of pore residues and ligands of the diiron(II) center in the hydroxylase-regulatory component complex of T4mo (PDB ID 3DHI).
Figure 4.
Substrate access through the cavities and the pore is triggered by binding of the regulatory component. a) In free hydroxylase of sMMO, cavities 1 and 2 are disconnected, and the pore is open (PDB ID 1MTY). Xe atoms were found in cavity 2 in MMOH crystals pressurized with Xe gas.\textsuperscript{44} b) When the regulatory component binds, the cavities are connected and the pore is closed (PDB ID 4GAM). The concerted motion of these two alterations is striking.
Figure 5. Channels in TMO. a) The channel in free hydroxylase of ToMO (PDB ID 2INC) and a magnified view of the channel in the inset. b) The channel in free hydroxylase of T4mo (PDB ID 3DHG). c) The channel in the hydroxylase-regulatory component complex of T4mo (PDB ID 3DHH). Channel in one of the hydroxylase monomer are shown; b) and c) only show one of the hydroxylase monomer. The channel and pore are shown in green and orange, respectively.