Toward Models for the Full Oxygen-Evolving Complex of Photosystem II by Ligand Coordination To Lower the Symmetry of the Mn₃CaO₄ Cubane: Demonstration That Electronic Effects Facilitate Binding of a Fifth Metal

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Supporting Information

ABSTRACT: Synthetic model compounds have been targeted to benchmark and better understand the electronic structure, geometry, spectroscopy, and reactivity of the oxygen-evolving complex (OEC) of photosystem II, a low-symmetry Mn₃CaO₄ cluster. Herein, low-symmetry MnIV-GdO₂ and MnIV-CaO₄ cubanes are synthesized in a rational, stepwise fashion through desymmetrization by ligand substitution, causing significant cubane distortions. As a result of increased electron richness and desymmetrization, a specific μ₃-oxo moiety of the Mn₃CaO₄ unit becomes more basic allowing for selective protonation. Coordination of a fifth metal ion, Ag⁺, to the same site gives a Mn₃AgCaO₄ cluster that models the topology of the OEC by displaying both a cubane motif and a “dangler” transition metal. The present synthetic strategy provides a rational roadmap for accessing more accurate models of the biological catalyst.

The oxygen-evolving complex (OEC) of photosystem II (PSII) is the sole biological water oxidation catalyst.¹ During the catalytic cycle, the OEC is photo-oxidized four times, affording the high-oxidation-state Mn necessary for O–O bond formation and release of O₂.² The structure of this heterometallic Mn/Ca cluster has been investigated through spectroscopic,² X-ray diffraction (XRD),³ and computational⁴ studies that support a low-symmetry Mn₃CaO₄ cluster containing a distorted Mn₃CaO₄ cubane bridged to a fourth manganese through one oxygen of the cubane and a μ₃-oxo. The redox-inactive Ca²⁺ has been shown experimentally to be crucial for catalytic activity,⁵ and multiple computational studies include the fourth, dangling Mn as an integral part of the catalytic cycle.⁴,⁶

Recently, the Mn₃CaO₄ cubane subsite of the OEC has been accurately modeled,⁷ providing electrochemical details suggesting that Ca²⁺ plays a role in tuning the reduction potentials of the Mn centers.⁷,⁸ Based on EPR and magnetism measurements, cluster asymmetry, or distortion, in the Mn₃CaO₄ unit, manifested in one case by coordination of a cubane oxo to a second Ca²⁺, was proposed to affect the cluster electronic structure and thus its chemical reactivity.⁷b Low-symmetry, heterometallic Mn₃MO₄ cubane complexes are therefore desirable synthetic targets for further electronic structure studies and as precursors to full Mn₃CaO₄ models of the OEC.

We have developed synthetic protocols to facilitate rational design of site-differentiated, homo- and heterometallic metal-oxo clusters.⁷a,⁸,⁹ The 1,3,5-triphenylbenzene-based ligand scaffold (L, Scheme 1) results in Mn₃MO₄ cubanes that have high, pseudo-C₃ symmetry and an apical metal, M, labile to substitution by more Lewis acidic ions.⁹ For example, Mn₃CaO₄ cubane complex 1 reacts quantitatively with Mnβ(OtTf)₂-2CH₃CN (OtTf = trifluoromethanesulfonate) to yield the reported Mn₃O₄ cubane 2 rapidly upon mixing (Scheme 1). This substitution prohibits appending a fourth Mn to 1 to generate a model of the full OEC. Herein, we report strategies to distort and stabilize Mn₃MO₄ cubanes, forming a pentametallic complex in the biologically relevant geometry as a first example of the Mn₃CaM₄O₄ unit with a dangling transition metal.

The Mn₃MO₄ cubanes supported by ligand L display alkoxide and pyridine coordination to the three MnO₂ faces (“bottom” of the cubane) and acetate bridging the three MnMO₂ faces (“top” of the cubane). Solvent molecules complete the coordination sphere of larger cations M (Scheme 1, complexes 1–3), such as Ca²⁺, Sc³⁺, Y³⁺, and Ln³⁺, with coordination numbers up to nine being observed.⁷a,⁸,⁹e Substitution of the solvent molecules and acetates with multidentate ligands offers the opportunity to rationally modify the structure and properties of the cubane. We selected the design elements of this multidentate ligand to address several aspects: (1) lower the propensity for substitution for the top metal, M, by a fifth metal equivalent, M′; (2) distort the cubane; and (3) change the reactivity of the cubane unit to allow incorporation of a fifth metal, M′. The MnIV-GdO₄ cubane 3 was chosen for initial studies since the size of Gd³⁺ affords more coordination sites and Gd³⁺ is less susceptible to metal substitution. The substitution of the acetates of MnIV-MO₄ cubanes is significantly slower than for cubanes displaying MnIII centers, as demonstrated by isotopic scrambling experiments with acetate-d₃ and acetate-d₄.⁹b Therefore, multidentate ligands were chosen that could bind not only to the top metal (Gd³⁺ or Ca²⁺) but also to at least one

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Mn$^{IV}$ of the cubane through substitution of acetates so that the new ligand would be stable to dissociation. Modification of the coordination sphere was achieved with N,N$'\text{-}$dimethyl-$N,N'$-diacetylene diamine dioxime (HON$_4$OH), which upon mixing with 3 and Et$_3$N in THF afford complex 4 in 90% yield. XRD studies show that the dioxime substitutes two of the acetates, resulting in two oximate bridged MnGdO$_2$ faces and four N-donors coordinated to Gd$^{3+}$. The Gd$^{3+}$ center is nine-coordinate with a DMF molecule completing the ligand sphere (Figure 1). The chelation of [ON$_4$O]$^{2-}$ to Gd$^{3+}$ is reminiscent of crown ether coordination to metals, generating multiple favorable five-membered rings. The binding of [ON$_4$O]$^{2-}$ generates a low-symmetry cubane core compared to 3. The structural parameters (Figure 2) show that the $\mu_3$-oxo moieties of 3 have more uniform Mn-O bond lengths (1.841(2)–1.916(2) Å), than in 4 (1.844(5)–1.946(4) Å), with the bonds trans to the oximate ligands elongated due to the higher trans influence of these donors compared to acetates. Additionally, the Gd-O2 distance is shortened, likely due to a combination of increased basicity at O2, increased acidity at Gd, and the chelating effect of the [ON$_4$O]$^{2-}$. The $^1$H NMR spectrum of 4 shows an increased number of peaks relative to 3 and the electrospray ionization mass spectrometry (ESI-MS) masses support bound [ON$_4$O]$^{2-}$ and loss of two acetates, both

“HON$_4$OH = N,N$'\text{-}$dimethyl-$N,N'$-diacetylene diamine dioxime. OTf$^-$ = triflate = trifluoromethanesulfonate.
Symmetry cubane structures could be built on ligand scaffolds, and the Mn$_3$CaO$_4$ system was targeted next as a more accurate model of the OEC. Addition of HON to Ca$_2^+$ instead of a proton at this position would be consistent with the low symmetry observed in the solid state persisting in solution.

With the Mn$_3$GdO$_4$ system as a proof-of-principle that low-symmetry cubane structures could be built on ligand scaffold L, the Mn$_3$CaO$_4$ system was targeted next as a more accurate model of the OEC. Addition of HON$_4$OH to LutHOTf in DCM (Supporting Information (SI), Figure S9). Therefore, the protonation of the Mn$_3$CaO$_4$ core specifically at O2 suggests that the [ON$_4$O]$_2^-$ ligand promotes (1) an increase in the electron richness of the cluster, as oximates are more basic than carboxylates, and (2) electronic desymmetrization of the cluster, together making the O2 site significantly more basic. This effect was further studied by quantum mechanics (B3LYP-D3 DFT, see SI) through the inspection of both canonical (nonlocal) and localized molecular orbitals. Ground-state optimizations of simplified models for 1 (1m), 5 (5m), and 6 (6m) were performed, showing good agreement with the crystal structures (RMS values of 0.024 Å (1) and 0.037 Å (6)). Analysis of localized molecular orbitals shows that the energy of O2 changes the most during ligation of [ON$_4$O]$_2^-$: The energies of the three equatorial μ-oxo motifs of 1m appear effectively degenerate (−0.72, −0.71, and −0.72 hartrees). Upon coordination of [ON$_4$O]$_2^-$, the energies of the oxides’ localized lone pairs are all higher in energy, consistent with a more electron rich cluster as expected for oximate vs acetate coordination. Notably, lowering the symmetry of the cubane (5m) leads to different lone pair energies for O2, O1, and O3 (−0.56, −0.69, and −0.66 hartrees), with the largest change occurring for O2. This asymmetry is reflected experimentally in the protonation of 5 at the O2 position to generate 6. Upon protonation of the most basic μ-oxo, the μ-hydroxo and the two μ-oxo moieties shift energy to −0.61 (σ$_{O^1−H}$ bond), −0.70, and −0.70 hartrees (lone pairs), respectively, in 6m. The canonical molecular orbitals reflect the notions above (SI, Figures S11–S13) by showing higher energy for the HOMO of 5m vs 1m and increase in electron density at the O2 position.

With the increased Lewis basicity of μ-oxo O2 established, binding of a “dangling” metal instead of a proton at this position was targeted to model the Mn$_3$CaO$_4$ + M’ geometry of the OEC.$^{12}$ Addition of metal triolate salts of Ag$_3^+$, Mn$_2^+$, and Co$_2^+$ to 5 led to changes in the $^3$H NMR and ESI-MS spectra, indicative of productive reactions (see SI). None showed loss of Ca$_2^+$ as compared to 1, a crucial finding for further cluster elaboration. Moreover, the ESI-MS spectra of the Mn and Co reactions displayed peaks consistent with LMn$_3$CaO$_4$(ON$_4$O)$^-$, LMn$_3$CaO$_4$(ON$_4$O)(OAc)$^-$, LMn$_3$CaO$_4$(ON$_4$O)(OAc)$^-$, LMn$_3$CaO$_4$(ON$_4$O)(OAc)$^-$, and LMn$_3$CaO$_4$(ON$_4$O)(OAc)$^-$, with fragmentation patterns corresponding to loss of OTf and H$_2$O, respectively, suggesting the formation of the desired precursor.

The symmetric Mn$_3$CaO$_4$ cubane 1 does not react with LutHOTf in DCM (Supporting Information (SI), Figure S9). The Mn$_3$CaO$_4$ core reveals that it is asymmetric. In particular, an outer sphere triolate anion is found in close proximity (O1T−O2 = 2.660(8) Å) to the μ$_3$-O moiety that coordinates to Ca$^{2+}$ and to the Mn centers supported by the oximates. This distance suggests protonation at this position to generate a μ$_3$-OH moiety involved in a hydrogen bonding interaction with the triolate.$^{10}$ In agreement, the Mn−O2 distances (1.926(5), 1.944(5) Å) of 6 are longer than in 1, 4 and the other top oxo O−Mn distances in 6 by 0.05−0.1 Å (Figure 2). Compound 6 is the first structurally characterized example of a protonated high oxidation state Ca−Mn mixed oxide cluster. The significant distortion promoted by protonation correlates to studies of the OEC.$^{11}$

Figure 3. Truncated solid-state structure of 6 and 7, with the core of 7 and the OEC overlaid.$^{2b}$ Hydrogens and outer-sphere anions not shown for clarity.
MnCaM’ stoichiometry in solution (SI, Figures S10–S15). Crystals suitable for XRD studies were obtained only from the reaction of S with AgOTf to generate cluster 7 (Figure 3). The quality of the XRD data is low, but does allow confirmation of connectivity. The solid-state structure of product 7 shows a Mn3CaAgO4 cluster. Due to the quality of the data the identity of the silver atom is not certain; however, the chemical coordination, bond distances, and displacement parameters are more consistent with Ag than Mn or Ca. The “dangling” Ag is directly linked to the cubane core via coordination to a μ-oxo, one μ-alkoxide, and one of the pyridines of L that was not coordinated in the precursors. The fourth coordination site is filled by a triflate. Ca2+ is in a similar geometry to 6. An overlay of the structures of the OEC and 7 shows that the geometry of the Mn3CaM’O4 clusters is well modeled (Figure 3). This represents the first example of rational, stepwise synthesis of the asymmetric pentametallic core model of the OEC. Starting from a trimanganese precursor, addition of Ca2+ and oxidation lead to the cubane core,7α which is then desymmetrized and further elaborated with addition of the “dangler” metal. Conceptually related, the photoreassembly of the OEC also involves several intermediates with stepwise incorporation of metals, although in a different order than demonstrated synthetically here.3,14 Additionally, the tuning of the basicity of coordinated OH moieties by metal ions has been invoked in the assembly of the biological cluster.15

In summary, starting from pseudo-C3 symmetric Mn33CaO4 and Mn33GdO4 cubanes, coordination of a multidentate bixoximate ligand leads to lower-symmetry clusters. Most notably, the oximate ligand stabilizes the Mn33CaO4 core against Ca2+ substitution. The cluster is affected electronically to increase the basicity of one of the oxo moieties, which can be selectively protonated to generate a Mn33CaO4(OH) cluster. The basicity of this site can also be exploited to coordinate a fifth metal. A Mn33CaAgO4 cluster was structurally characterized, showing high structural similarity to the OEC. The present compounds provide insight into the electronic tunability of clusters related to the OEC by ligand binding. Additionally, the reported strategy provides the first rational synthetic roadmap for accessing more accurate models of the biological catalyst.

■ ASSOCIATED CONTENT

Supporting Information
Synthetic procedures, spectroscopic characterization, crystallographic data, and computational details and results. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
The authors declare no competing financial interest.

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