Oxygen Production in Nature: A Light-Driven Metalloradical Enzyme Process

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Introduction

Dioxygen is thermodynamically hot but kinetically cool, which makes it an ideal reagent for maximizing biological free energy production and for carrying out difficult chemical transformations in enzyme active sites. The widespread use of dioxygen in biological catalysis has led to an enzyme classification scheme — monooxygenases, dioxygenases, oxidases — that is based on the specifics of the chemistry in which O2 participates. Examples of the remarkable utility of dioxygen in biology abound and include its use in maximizing ATP production in aerobic respiration, in C–H bond activation in the P450 enzymes and methane monooxygenases, and in the degradation of important biomaterials such as lignin.

Although nature has devised a multitude of mechanisms by which to activate dioxygen for useful chemistry, only one system, Photosystem II (PSII) in plants and algae, has evolved that has the capacity to lift water out of its thermodynamic well to generate dioxygen. This singular system, Photosystem II, provides the substrate-binding sites in the water-splitting complex. Recent work has shown that the tyrosine, the proline, and Cl– that accompany oxygen reduction to water in oxygen-metabolizing metalloproteins are likely to function as a de novo protein synthesis.

Each photon-absorption and charge-separation event in PSII provides only a single oxidizing equivalent to the OEC of PSII, as waste is coming into view. In this article, we review the physical structure and energetics of PSII. We then discuss and analyze a metalloradical enzyme mechanism for the water-oxidation process it catalyzes.

Physical Structure of Photosystem II and an Overview of Water Oxidation

PSII occurs as a multi-subunit, membrane-spanning complex in the thylakoid membrane. Figure 1 shows a scheme of its reaction center that includes a number of the polypeptides and redox cofactors that are essential to the photochemistry and subsequent electron- and proton-transfer reactions that precede oxygen evolution. Light absorption by the specialized chlorophyll complex, P680, generates a charge-separated state, P680+QA–, that is stable to recombination for ~200 μs. Reduction of P680+ by a redox-active tyrosine YZ, Y161 of the D1 polypeptide, is fast and outcompetes the wasteful charge-recombination process. Electron transfer from QA to the secondary acceptor, QB, and concomitant protonation mobilize the hydroquinone for electron delivery to Photosystem I and, eventually, the CO2-fixing reactions of the Calvin cycle.

YZ deprotonates as it reduces P680+ to produce the neutral YZ radical. Reduction and reprotonation of YZ occur from water and the tetranuclear (Mn)4 complex that provides the substrate-binding sites in the water-splitting complex. Recent work has shown that the tyrosine, the (Mn)4 cluster, and the other two cofactors essential for water oxidation, Ca2+ and Cl–, are likely to function as a single catalytic center in forming the oxygen-evolving complex (OEC) of PSII, as discussed below.

FIGURE 1. Major polypeptides and redox components in PSII.
by postulating the S-state notation in which the number of oxidizing equivalents stored in the OEC is denoted as $S_n$. The photochemistry/oxidizing equivalent accumulation process can thus be described as follows:

$$S_n P_680 Q_A^+ Q_B^- \rightarrow S_{n+1} P_680 Q_A^+ Q_B^-$$

Only when the $S_4$ state is reached does water oxidation occur.

**Energetics in PSII Photochemistry and Water Oxidation**

There are stringent energetic constraints on PSII that provide insight into viable mechanisms for water oxidation. Figure 2 compares the photochemical energetics of PSII with that of reaction centers from photosynthetic purple bacteria. The latter, for which detailed crystallographic information is available, has structural and functional analogies to PSII and is often used as a template in discussing PSII operation. This approach has clear predictive power, as demonstrated by similarities in the quinone regions of the two photosystems and in the extension of $C_2$ symmetry themes to PSII.

Nonetheless, there are key differences. As Figure 2 shows, the efficiency of the conversion of photon energy into chemical potential is only 35% in the bacterial system, whereas it approaches 70% in PSII. This disparity arises from differences in the extent to which energy transfer and redox transitions are thermodynamically downhill in the two systems. The bacterial reactions are strongly driven, and the free energy changes enroute to a stable charge separation are large. An absorbed photon leads relentlessly to charge separation but with a considerable sacrifice in the chemical potential that is developed in the charge-separated state.

In PSII, weaker coupling occurs between chromophores and between redox cofactors, and the degree of irreversibility in each step lies less strongly in favor of product.

**Construction of the Catalytic Center for O₂ Evolution**

Figure 4 provides a model for the oxygen-evolving complex that combines recent data in the literature with charge neutrality and proton current considerations. Consensus that the manganese ions required for water splitting occur as a compact tetranuclear cluster is emerging. Yachandra et al. have interpreted their X-ray absorption data to indicate that the (Mn)₄⁺ center has a C shape in their structure, one can calculate that a distance of 5.5 Å separates the two manganese ions at the open end of the C. Although more controversial, the idea
that the Ca$^{2+}$ and Cl$^-$ cofactors are closely associated with the cluster has reasonable support.2b,9a,10

Site-directed mutagenesis has identified potential ligands to the (Mn)$_4$/Ca$^{2+}$ cluster, notably D170, H332, E333, H337, D342, and the carboxy terminus of A344, all in the D1 subunit of PSII.3,11 YZ is close to the metal cluster,4,12,13 and the proton-accepting base, which allows the functionally critical deprotonation of YZ during its oxidation by P680$^+$, is probably D1 H190.3 To account for the appearance of protons in bulk phase on the time scale of YZ oxidation,14 we suggested that H190 communicates with the inner thylakoid aqueous space by a hydrogen-bonded chain that may involve D1 E189,3 either directly or by positioning water molecules in a hydrogen-bonded network analogous to those for the quinone sites in the bacterial reaction center15 and for the binuclear center in cytochrome oxidase.16

In S$_0$, the valences of the four Mn ions sum to 13+.2b,9a We assign these to individual ions as 4+, 4+, 3+, and 2+; including the Ca$^{2+}$, the overall positive charge on the cluster in S$_0$ is 15+. This positive charge is exactly balanced by the five bridging oxos, the four carboxylates noted above, and the Cl$^-$. Moreover, these ligands are arranged in Figure 4 to provide local as well as overall charge neutrality in the cluster, i.e., so that each of the five metals has zero net charge. Electro neutrality in protein-bound metal centers is general17 and follows from the energetic consequence of attempting to bury charge in regions of low dielectric.18 The structures of many metalloprotein active sites, for example, that of the binuclear Mn cluster in arginase19 and of the binuclear Fe centers in ribonucleotide reductase and methane monoxygenase,20 demonstrate electroneutrality and its persistence during redox changes at the metal centers. Charge can apparently be tolerated on a metal/ligand cluster only when the site is sufficiently close to the surface of the protein to experience significant electrostatic screening from solvent water.17 As the OEC is remote from bulk phase,21 solvent screening will be muted, which reinforces the likelihood of the electroneutral structure shown in Figure 4. The operation of the metal-cluster/YZ site in oxygen evolution, described below, is driven to a large extent by the necessity of preserving overall electroneutrality in the complex during the S-state cycle.5

The Role of YZ in Water Oxidation and the S-State Cycle

YZ reduces P680$^+$ in nanoseconds to preserve the high quantum yield of charge separation in PSII and the highly oxidizing potential associated with P680$^+$. Recent work has extended the role of the tyrosine in PSII and implicated YZ$^-$ intimately in the water-oxidizing chemistry itself.4,5 By combining experimental data on YZ$^-$ with emerging ideas on the function of the radical moiety in the general class of metalloradical enzymes, we suggested that YZ carries out hydrogen-atom abstraction from substrate water on each S-state transition.5,22

The rationale for this model is summarized in Figure 5. Figure 5A shows a mechanistic scheme for O$_2$-depend-

![Figure 5](https://example.com/figure5.png)

FIGURE 5. Schematic for casting PSII as a member of the metalloradical enzymes class.

ent metalloradical enzymes.23 In this class of proteins, which includes galactose oxidase, ribonucleotide reductase, and prostaglandin synthase, the metal site activates dioxygen to produce a strong metal-bound oxidant. This activated oxygen species generates the amino acid radical by H-atom abstraction and the radical, in turn, abstracts a hydrogen atom from substrate to initiate catalysis. In casting PSII as a member of this class, we reverse the direction of the H-atom current (Figure 5B) and recognize that P680$^+$ is sufficiently oxidizing to serve as the thermodynamic sink in this process (Figure 5C). By postulating this mode of operation for PSII, clear analogies between it and other metalloradical enzymes emerge.5,22

The mechanistic view of PSII in Figure 5 meshes well with the structural model in Figure 4. The tyrosine is ideally positioned to promote forward electron transfer to P680$^+$, which is located 7–12 Å from it,20 while simultaneously undergoing deprotonation through the H190/E189/.../bulk phase pathway. This structure emphasizes the tight coupling in PSII between electron- and proton-transfer events that Krishtalik's work6 virtually insists upon (see below). The proximity of YZ and the metal cluster is ideal for the H-abstraction function postulated for YZ$^-$.

We identify the two Mn ions at the open end of the C as the catalytic ions that bind substrate; H-atom abstraction by YZ$^-$ from water bound terminally at these catalytic sites lies at the heart of each of the S-state transitions. Thus, the function of manganese in water oxidation becomes clear: it serves to bind substrate water, to delocalize oxidizing equivalents that are generated upon each H-atom abstraction, and to provide a template for
the critical O–O bond formation step. For the latter process, the 5.5 Å separation between the two catalytic ions is ideal.

Figure 6 presents a full S-state model based on the H-atom abstraction model. Although necessarily speculative in some aspects, it is consistent with a wide variety of observations on O₂ evolution. In S₀, the catalytic sites have 2⁺ or 3⁺ valences, and each is occupied by substrate water (see below). Photogeneration of P680⁺ initiates the electron and proton motions that produce Y₂⁺. The radical, in turn, abstracts an H atom from water bound to the upper catalytic ion to produce a ligated hydroxide; the oxidizing equivalent delocalizes to the Mn to oxidize it to 3⁺ in S₁. Upon a second photon absorption, Y₂⁺ is formed and abstracts an H atom from the hydroxide to produce, transiently, an oxo species in the state labeled S₂*. For the d⁴ Mn³⁺ ion in the S₁ state, Jahn–Teller (JT) distortions are in effect and coordination to the axial position is expected to be weak or absent. Upon oxidation to the d³ Mn⁴⁺ state in S₂*, the JT distortion is lifted. We postulate that relief of this distortion provides a driving force for Cl⁻ motion to the upper catalytic site. Ligation of the anion to the Mn⁴⁺ in S₂* increases the basicity of the oxo, relative to the water bound to the lower catalytic ion, which promotes a proton-transfer process to produce hydroxides at both catalytic centers. The coupled H⁺/Cl⁻ motion is essential in positioning substrate in the higher S states and completes the S₂ → S₃ transition in Figure 6. Anion ligation may also prevent premature water oxidation. Upon atom abstraction in S₂ → S₃, the hydrogen-bonded oxo/hydroxyl species is formed. These ligands may equilibrate with a protonated peroxy species (Mn⁴⁺=O–HO–Mn⁴⁺), although the equilibrium cannot lie too far right, as net manganese reduction does not take place on S₂ → S₃. The final, O₂-producing transition proceeds according to the atom-abstraction/radical addition process discussed elsewhere. Upon completion of the cycle, Cl⁻ migrates back to the Ca site and two water molecules bind to reform the S₀ state.

The model in Figure 6 is structurally and mechanistically well founded. Moreover, it maintains overall cluster electroneutrality throughout the S-state cycle, which avoids the significant energetic penalties that would occur if charge were allowed to accumulate. The model is conservative, as essentially the same chemistry occurs on each S-state transition, and minimal nuclear motion is required within the complex as it cycles through to produce O₂. The latter feature is of considerable importance, both in terms of the low driving force available to split water in PSII and of the high turnover number (up to 200 electrons s⁻¹) of the OEC.

Thermodynamics and Kinetics of the H-Atom Abstraction Process

The mechanism in Figure 6 must be shown to be thermodynamically feasible and kinetically competent. At first glance, the abstraction process presents a thermodynamic problem, as the bond dissociation energies (BDE’s) for water (119 kcal/mol) and hydroxyl (102 kcal/mol) are significantly more exothermic than that for the tyrosyl O–H bond (87 kcal/mol). Figure 7 summarizes the uncatalyzed situation. The formation of O₂ with a tyrosyl radical (Y(·O)) is exothermic as required, but only because of the strong exothermicity for forming the O=O bond in the final step; the individual H-abstraction steps are strongly endothermic.

\[
\frac{1}{2} _2H_2O + Y(·O) → \frac{1}{2} _2O_2 + Y(OH)
\]
Oxygen Production in Nature  Tommos and Babcock

FIGURE 7. Comparison of the BDE’s for water, free in solution$^{28b}$ \((\text{Y}^+ + \text{H}_2\text{O})\) (catalyzed), and when complexed by a manganese ion$^{31}$ \((\text{Y}^+ + \text{H}_2\text{O} \text{(catalyzed)})\). The horizontal line represents the BDE for the tyrosine O–H bond.$^{28a}$ The vertical arrows show BDE differences between substrate water or hydroxyl O–H bonds and the tyrosyl O–H. The upward arrows, for the uncatalyzed case, indicate that atom abstraction by the tyrosyl radical is endothermic; the downward arrows, for the catalyzed case, indicate the exothermicity of abstraction by the tyrosyl radical. Two cycles, i.e., the process for two water molecules, are shown to demonstrate the energetics of \(\text{O}_2\)-bond formation as the right-most downward arrows.

But, these energetics change radically when water is ligated to manganese.$^{29-31}$ The BDE’s for H-atom abstraction drop for both water and hydroxide and occur in the 77–87 kcal/mol range (Figure 7), consistent with the essentially thermoneutral character of the \(\text{Y}_Z\text{S}_n\) transitions. The individual S-state transitions are now thermodynamically appropriate, and the O–O bond forming step is less strongly driven.

In radical-driven atom abstraction processes, entropic factors are close to zero,$^{32}$ which supports Mayer’s emphasis on the predictive utility of BDE differences in radical reactions.$^{29}$ The importance of the energy release during the \(\text{Y}_Z\text{O}–\text{H}\) bond forming process and the nulling of entropic effects dovetail well with Krishtalik’s analysis of \(\text{O}_2\) evolution.$^8$ He noted that the configurational potential for the four-electron water-oxidation process

\[
\frac{1}{2}\text{H}_2\text{O} = \frac{1}{4}\text{O}_2 + \text{H}^+ + \text{e}^{-} \tag{2}
\]

is 1.401 eV, which is significantly greater than the driving potential of 1.12 eV available from the reduction of P680$^+$. Accordingly, he concluded that water oxidation in PSII, in the absence of additional favorable processes, is not kinetically competent.

This realization led him to examine ways by which the OEC coupled further reactions to the fundamental redox chemistry in reaction 2. He quickly eliminated two such possibilities by showing that binding the oxidized oxygen-containing product species or beginning the reaction with hydroxide rather than water is ineffectual. However, protonation of a base (X$^-$) as the redox chemistry in (2) occurs, i.e.,

\[
\frac{1}{2}\text{H}_2\text{O} + \text{X}^- = \frac{1}{4}\text{O}_2 + \text{XH} + \text{e}^{-} \tag{3}
\]

is more fruitful. His analysis showed that proton-binding affects the configurational potential as follows:

\[
E'_c = E_c + \frac{1}{2}E_{\text{H}_2\text{O}} - [0.059(pK_a - \log X_{\text{H}^+})] \tag{4}
\]

where \(E'_c\) is the configurational potential for (2) (1.401 V), \(E_{\text{H}_2\text{O}}\) is the binding energy of \(\text{H}_2\text{O}\) in the OEC, \(pK_a\) is that for the base in (3), and \(\log X_{\text{H}^+}\) is the mole fraction of protons at pH 0 (1/55.5). With a reasonable binding energy for water (0.3 eV), a graphical analysis of eq 4 shows that \(E'_c\) drops to values below that of \(\text{Y}_Z/\text{Y}_Z\) for \(pK_a\)'s for \(\text{XH}^+\) greater than 7.5 (Figure 8).

If the \(pK_a\) of \(\text{XH}^+\) in (3) approaches that of bulk solution, however, it is substantially protonated prior to the electron transfer and, thus, unavailable as a proton acceptor. If the \(pK_a\) is decreased to overcome this, the effect of the coupled protonation on the configurational potential is diminished and \(E'_c\) rises. To circumvent this difficulty, Krishtalik suggested redox-linked dissociation of a manganese ligand, possibly a semiquinone, to provide an acid/base function whose \(pK_a\) was coupled to the electron transfer in (2). This mechanism now appears unlikely, as quinones are not associated with the OEC, and there are no data to support the idea of redox-linked ligand dissociation on each S-state transition.

The tyrosyl radical in Figures 4 and 6, however, fulfills Krishtalik’s criteria precisely. Upon oxidation by P680$^+$, the tyrosyl proton is sloughed from the site, owing to the low \(pK_a\) (≈2) of the cation radical, to generate \(\text{Y}_Z\). Its subsequent rereduction by H-atom transfer from substrate water/hydroxide regenerates the tyrosine form with \(pK_a\) ~ 9. Within the context of (3), this \(pK_a\) is the operational value. We can represent the reaction sequence involved as

\[
\text{Y}_Z^+ + \text{e}^- \leftrightarrow \text{Y}_Z^- \tag{a}
\]

\[
\frac{1}{2}\text{H}_2\text{O} + \text{Y}_Z^- \leftrightarrow \frac{1}{4}\text{O}_2 + \text{Y}_Z + \text{e}^- \tag{b}
\]

\[
\frac{1}{2}\text{H}_2\text{O} + \text{Y}_Z^+ \leftrightarrow \frac{1}{4}\text{O}_2 + \text{Y}_Z \tag{net} \tag{5}
\]

where \(\text{Y}_Z\) represents the reduced, protonated tyrosine species. The high \(pK_a\) for \(\text{Y}_Z/\text{Y}_Z^+\) reduces the configurational potential for step b to ~0.9 V (Figure 8), and the
and K

where H), they developed the LFE

For phenoxyls reacting with organic hydroperoxides (ROO

the weak driving forces available. 2 Can the metalloradi-

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Figure 9 summarizes these LFE’s graphically. Ingold and

co-workers recently presented a study of phenoxyl radical

reactivity consistent with the above.33

FIGURE 9. Linear free-energy relationships for H-atom abstraction

processes according to eqs 6–8. The shaded area represents the

range of the equilibrium constants for the S,Y2* ↔ S,Y2 reactions of

the S-state cycle.2

configurational potential available in the reduction of Y2*

(−1.0 V) becomes sufficient to drive the net process in

(5). Thus, the Krishtalik analysis within the context of the

metalloradical mechanism for water oxidation (5) con-

verges with the BDE data for reaction 1.

The S-state transitions have rates of 103−104 s−1, despite

the weak driving forces available. 2 Can the metalloradi-

cal—enzyme mechanism account for these kinetics? For

atom-abstraction processes, linear free energy relations-

ships (LFE’s) that relate the difference in BDE’s for the

reactant and product bonds (ΔBDE) to the rate of the

abstraction process are available. Mahoney and Da-

rooge,32 for example, have studied the relationship be-

between rate and equilibrium for H-atom abstraction from

both O−H and C−H bonds by phenoxyl radicals (PhO*).

For phenoxyls reacting with organic hydroperoxides (ROO−

H), they developed the LFE

\[ \log k = 4.0 + 0.51 \log K_{eq} \]  

(6)

where k is the second-order rate constant for abstraction

and K_{eq} is the equilibrium constant for the abstraction

process. For the analogous reactions of phenoxyl radicals

with substituted phenols (RO−H), the abstraction rate was

independent of K_{eq}

\[ \log k = 5.65 + 0.15 \]  

(7)

where k is defined as in (6). Finally, for phenoxyl radicals

abstracting H atoms from hydrocarbon C−H bonds, the

relationship

\[ \log k = -1.2 + 0.2 \log K_{eq} \]  

(8)

was observed, where k and K_{eq} are defined as for (6). Figure 9 summarizes these LFE’s graphically. Ingold and

co-workers recently presented a study of phenoxyl radical reactivity consistent with the above.33

These LFE’s led to two important conclusions. First,

phenoxyl radicals are significantly, by 4–5 orders of

magnitude, more reactive in abstracting H atoms from

O−H bonds than from C−H bonds. Thus, assessment of

the kinetic competence of the mechanism in Figure 6 on

the basis of LFE’s derived from C−H bond hydrogen-atom

abstraction rates is not well founded. Second, if we take

the effective concentration of the reacting partners in the

OEC, Y2 and Mn^{+} H_{2}O(OH), as 10 M^{2b} and equilibrium

constants in the range 2−10 for the individual S-state transitions, 2 rate constants for these transitions of \( \geq 10^{6} \)

s−1 are predicted (Figure 9). The lower rates, 103−104 s−1, observed for S,Y2* → S,Y1+Y2 may reflect metal ligation or

protein constraints. Overall, the organic radical reactivity
data show that Y2 is kinetically competent to carry out the

H-atom abstraction processes we postulate.

The Y2* species could provide a hazard in PSI, as the

BDE’s of various protein C−H bonds are close to the

tyrosyl O−H bond.28,34 Thus, destructive side reactions

become possible. The number of nonexchangeable pro-
tons in the vicinity of the radical, however, is low in Mn-

containing PSI preparations, which suggests that the
catalytic site is protected from spurious reactions by

isolating Y2* from C−H bonds with which it could react.35

Electroneutrality and Substrate Binding

Britt reviewed work aimed at understanding the steps in

the S-state cycle at which substrate water/hydroxide binds,26

and concluded that water or hydroxide is already ligated in the lower S states. This conclusion is consistent

with recent ENDOR measurements36 and suggests that two

substrate molecules reload directly following O2 formation

and are bound at the catalytic Mn sites in S0 (Figure 6).

Pecoraro and co-workers studied the acid/base properties

of Mn model compounds.30,37 This work, in combination

with the likelihood of electroneutrality in the OEC, provides insight as to the protonation state of the bound

waters, i.e., whether substrate ligates as water or hydrox-

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and mononuclear Mn complexes. From this compilation,

several trends emerge. First, overall cluster charge is

critical in determining acid/base properties. Introducing

a charge as the result of deprotonation, whether at

bridging (compounds 1 and 5) or terminal (9) positions,

requires extremely basic conditions (pK_{a}’s 19–25), and the

oxos become progressively more acidic as the total charge

in the cluster increases. Second, the manganese valence,

within the III and IV valence set that is most relevant to the

OEC, has little effect on p


charge of 1

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The oxygen-evolution chemistry in PSII arises from a remarkable enzymatic structure that operates under stringent energetic and kinetic constraints. We have set out many of these and have described a mechanism that operates within the confines they impose. The mechanism survives thermodynamic and kinetic tests, as described above, and provides a chemical basis for the formation of the $O=O$ bond. A specific structure for the (Mn)$_4$ cluster has been adopted in developing this mechanism, and we have proposed roles for Ca$^{2+}$ and Cl$^{-}$ within the context of hydrogen-atom abstraction chemistry. These aspects of the model are based on solid data from a variety of labs, but they continue to attract intense experimental scrutiny; the details of these facets of Figure 6 may evolve. Nonetheless, the concepts that underlie the mechanism — $H$-atom abstraction on each S-state transition, the importance of overall charge neutrality, and the functional association of $Y_{Z}$, (Mn)$_4$, Ca$^{2+}$, and Cl$^{-}$ in forming a structurally compact water-splitting complex — are likely to persist and provide a solid basis for further effort to understand the chemistry that pumps $O_2$ into our atmosphere.

**Conclusions**

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**Table 1. pK_a's of Selected Dimanganese Compounds**

<table>
<thead>
<tr>
<th>no.</th>
<th>compd</th>
<th>$\Delta q$</th>
<th>pK_a</th>
<th>ref</th>
</tr>
</thead>
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<tr>
<td>1</td>
<td>[Mn$^{IV}$Mn$^{III}$ (salpn)$_2$]$^f$</td>
<td>0 → 1</td>
<td>24.5 ± 0.7$^c$</td>
<td>30a</td>
</tr>
<tr>
<td>2</td>
<td>[Mn$^{IV}$Mn$^{IV}$ (salpn)$_2$]</td>
<td>1 → 0</td>
<td>13.4 ± 0.2$^c$</td>
<td>30a</td>
</tr>
<tr>
<td>3</td>
<td>[Mn$^{IV}$Mn$^{III}$ (salpn)$_2$]</td>
<td>1 → 0</td>
<td>13.0$^f$</td>
<td>30a</td>
</tr>
<tr>
<td>4</td>
<td>[Mn$^{IV}$Mn$^{III}$ (salpn)$_2$]</td>
<td>2 → 1</td>
<td>6.5$^f$</td>
<td>30a</td>
</tr>
<tr>
<td>5</td>
<td>[Mn$^{IV}$Mn$^{III}$ (3,5-dCl (salpn)$_2$]</td>
<td>0 → 1</td>
<td>20.5 ± 1.0$^f$</td>
<td>30a</td>
</tr>
<tr>
<td>6</td>
<td>[Mn$^{IV}$Mn$^{III}$ (3,5-diCl (salpn)$_2$]</td>
<td>1 → 0</td>
<td>10.8 ± 0.3$^f$</td>
<td>30a</td>
</tr>
<tr>
<td>7</td>
<td>[Mn$^{III}$Mn$^{III}$ (bpy)$_2$]</td>
<td>2 → 3</td>
<td>11.0$^f$</td>
<td>37</td>
</tr>
<tr>
<td>8</td>
<td>[Mn$^{IV}$Mn$^{III}$ (bpy)$_2$]$^f$</td>
<td>4 → 3</td>
<td>2.3$^f$</td>
<td>37</td>
</tr>
</tbody>
</table>

$^a$ The $\Delta q$ is the net change in charge on the complex as it undergoes deprotonation. $^b$ salpn = N,N'-bis(salicylidene)-1,3-propanediamine. $^c$ Measured in CH$_3$CN or corrected to reflect pH. $^d$ Measured in CH$_3$CN or corrected to reflect pK_a. $^f$ Measured in H$_2$O. $^g$ Formed by P680. $^h$ L = 2-hydroxy-1,3-bis(salicylamino)propane. $^i$ Measured in 16 M water/CH$_3$CN.

in the S$_0$ state, which is unusual in protein-bound clusters, would we expect to ligate hydroxide in effective competition with water. Thus, the model compound work provides a strong basis for H$_2$O as the ligated substrate form.

**Calcium, Chloride, and Proton Currents**

The H-atom abstraction model predicts proton release upon each S-state transition with release kinetics on the time scale of Y$_Z$ oxidation. This behavior is observed in a variety of thylakoid, PSII-membrane, and PSII-core preparations. Moreover, in PSII-core particles, which are taken to reflect a combination of substrate proton release pattern. The proton-release pattern is more intricate, which is usually taken to reflect a combination of substrate proton release pattern and S-state-dependent redox Bohr effects. In addition, a persistent electrochromism accompanies the S$_1$ → S$_2$ transition, which is reversed on the S$_2$ → S$_3$ → S$_0$ transition. These band shifts are not coupled to the proton-release pattern. Consistent with the arguments above on electroneutrality, we attribute both the persistent electrochromic effects and the Bohr effects on H$^+$ release to internal charge rearrangement within the OEC, rather than to the creation of a naked charge.

The S-state cycle in Figure 6 accounts for this internal charge rearrangement. In S$_0$ and S$_2$, local electroneutrality is maintained about each of the metal centers in the OEC. However, with the generation of the transient oxo species in S$_2^*$, we postulate coupled H$^+$ and Cl$^-$ motion to form an S$_0$ state competent to complete the S cycle. In S$_2$, local neutrality is maintained about the upper catalytic Mn ion (two bridging oxos, the Cl$^-$ and the OH$^-$), but a H$^+$/H$_2$ dipole is introduced between the lower catalytic Mn and the Ca. We attribute the local electromorphism and the Bohr effects to this internal charge rearrangement. Cl$^-$ motion is clearly implicated in these phenomena, as neither is observed on the S$_0^-$ → S$_2^-$ transition in Cl$^-$-depleted material, even though Mn oxidation occurs. Moreover, the charge rearrangement upon S$_2$ formation may rationalize the decrease in the rate of Y$_Z$ oxidation by P680$^+$ in the higher S states.

By postulating a mechanism involving Cl$^-$ in positioning substrate for H-atom abstraction on S$_2$ → S$_1$ and S$_3$ → S$_0$. Figure 6 accounts for a variety of observations on the roles of Cl$^-$ and Ca$^{2+}$. Thus, Ca$^{2+}$ serves as an anchor for Cl$^-$ within the catalytic site, consistent with the ordered binding of these two cofactors and the slow CI$^-$ release kinetics. The 17 and 24 kDa polypeptides play a role in the latter phenomena and, together, are likely to prevent Cl$^-$ association/dissociation from rate-limiting water oxidation. Under conditions of Cl$^-$ or Ca$^{2+}$ depletion, the S-state cycle can advance one step beyond the S$_0$ state; subsequent illumination generates the modified S$_3$ transition in Cl$^-$-deficient material. The linked action of Ca$^{2+}$ and Cl$^-$ in controlling proton motion for the formation of the functional S$_2$ state provides a rationale for the susceptibility of the oxygen-evolving complex to produce this modified S$_2$ state, which cannot advance further in the S-state cycle, under a variety of inhibitory conditions.

**Oxygen Production in Nature**

Tommas and Babcock
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References


