Living with Oxygen
Harry B. Gray*Ω and Jay R. Winkler*Ω

Beckman Institute, California Institute of Technology, Pasadena, California 91125, United States

ABSTRACT: Work on the electronic structures of metal–oxo complexes began in Copenhagen over 50 years ago. This work led to the prediction that tetragonal multiply bonded transition metal–oxos would not be stable beyond the iron–ruthenium–osmium oxo wall in the periodic table and that triply bonded metal–oxos could not be protonated, even in the strongest Bronsted acids. In this theory, only double bonded metal–oxos could attract protons, with basicities being a function of the electron donating ability of ancillary ligands. Such correlations of electronic structure with reactivity have gained importance in recent years, most notably owing to the widespread recognition that high-valent iron–oxos are intermediates in biological reactions critical to life on Earth.

In this Account, we focus attention on the oxygenations of inert organic substrates by cytochromes P450, as these reactions involve multiply bonded iron–oxos. We emphasize that P450 iron–oxos are strong oxidants, so strong that they would destroy nearby amino acids if substrates are not oxygenated rapidly; it is our view that these high-valent iron–oxos are such dangerous reactive oxygen species that Nature surely found ways to disable them. Looking more deeply into this matter, mainly by examining many thousands of structures in the Protein Data Bank, we have found that P450s and other enzymes that require oxygen for function have chains of tyrosines and tryptophans that extend from active-site regions to protein surfaces. Tyrosines are near the heme active sites in bacterial P450s, whereas tryptophan is closest in most human enzymes. High-valent iron–oxo survival times taken from hole hopping maps range from a few nanoseconds to milliseconds, depending on the distance of the closest Trp or Tyr residue to the heme. In our proposed mechanism, multistep hole tunneling (hopping) through Tyr/Trp chains guides the damaging oxidizing hole to the protein surface, where it can be quenched by soluble protein or small molecule reductants. As the Earth’s oxygenic atmosphere is believed to have developed about 2.5 billion years ago, the increase in occurrence frequency of tyrosine and tryptophan since the last universal evolutionary ancestor may be in part a consequence of enzyme protective functions that developed to cope with the environmental toxin, O2.

INTRODUCTION

Human life on our planet could not exist were it not for two redox reactions: water oxidation to oxygen in photosynthesis and oxygen reduction to water in respiration. Also of enormous importance is the oxygenation of organic molecules by cytochrome P450. Years of research have shown that key reaction intermediates in these processes are multiply bonded oxo complexes of high-valent metals: cysteine-bound heme iron–oxos (compound 1) in P450,1 a manganese–oxo (or oxyl) in the oxygen evolving complex (OEC) of photosystem II,2 and a histidine-bound heme iron–oxo in cytochrome c oxidase, the terminal enzyme in respiration.3 As it also is known that these metal–oxos are very powerful oxidants, we might expect that living cells full of organic molecules would suffer from oxidative destruction. The million dollar question is then: how do we live happily on a planet bathed in oxygen?

Our story started in Copenhagen, nearly 60 years ago, when one of us (H.B.G.) began an investigation of metal–oxo bonding. Employing a modified Mulliken molecular orbital (MO) theory for calculations of orbital energies, the d1 vanadyl ion was shown to possess a V=O–oxo triple bond (V≡O), with a dxz,dyz(π*) level well above a singly occupied dx2–y2 (nonbonding) orbital.4 In this axially compressed tetragonal ligand field, the dipole-allowed dπ → dxz,dyz(π*) absorption accounted for the blue color of the triply bonded oxo ion. Related work published soon after predicted that absorptions attributable to analogous low energy transitions would be observed in all multiply bonded (d1–d3) metal–oxo complexes.5

THE OXO WALL

The first two electrons in a C4v M(O)L5 complex (Figure 1) will occupy the dπ orbital: the π-bond order will be 2 in d2, d1, and d0 cases, and it will decrease to 3/2, 1, and 1/2 for d3,d 4, and d5 complexes, respectively, as energetic dπ,dπ* π* orbitals are populated.6 It follows that d0 metal–oxo bonds will be considerably weaker [ν(M–O) ≈ 800 cm−1]6 than those of d1–2 mono-oxos [ν(M–O) ≈ 900–1000 cm−1].7,8 Three π* electrons cannot be tolerated, so multiply bonded tetragonal d3 oxos are not stable. With a d0 limit, then, an oxo wall separates groups Fe–Ru–Os and Co–Rh–Ir in the periodic table (Figure 2).9

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oxo multiple bonding, leading to a shift of the wall to the right of orbitals (Figure 3). In complexes with 1 or 2 d electrons, the oxo bond length, and a reduction in the energy gap between least one of those electrons must populate e\textsuperscript{π} orbitals and the M–oxo π-antibonding e\textsubscript{π} (d\textsubscript{xy},d\textsubscript{yz}) orbital.

Terminal oxo ligand leads to a large energy gap (>10,000 cm\textsuperscript{-1}) between the nonbonding b\textsubscript{2}(d\textsubscript{xy}) orbital and the M–oxo π-antibonding e\textsubscript{π} (d\textsubscript{xy},d\textsubscript{yz}) orbital. Strong \( \pi \) (superscript \( \pi \ )) coordinate linear (or 5-coordinate trigonal (C\textsubscript{2v})), 3-coordinate planar (C\textsubscript{3v}), or 2-coordinate linear (C\textsubscript{1v}) complexes, d\textsuperscript{6} configurations can support M–oxo multiple bonding, leading to a shift of the wall to the right (orange line).

**Metal–Oxo pK\textsubscript{a} Values**

In the tetragonal metal–oxo complexes described by the molecular orbital model depicted in Figure 1, the M–oxo bond order and, hence, the pK\textsubscript{a} of the conjugate acid (MOH\textsuperscript{+}) are sensitive functions of the population of the e\textsubscript{π}\textsuperscript{π} (d\textsubscript{xy},d\textsubscript{yz}) pair of orbitals (Figure 3). In complexes with 1 or 2 d electrons, the M–oxo π-bond order is two. In d\textsuperscript{3} configurations, however, at least one of those electrons must populate e\textsubscript{π}\textsuperscript{π} (d\textsubscript{xy},d\textsubscript{yz}), leading to a reduction in M–oxo π-bond order, an increase in the M–oxo bond length, and a reduction in the energy gap between e\textsubscript{π}\textsuperscript{π} and \( b_2 \) (\( \Delta E_{p} \)). The reduction in \( \Delta E_{p} \) as electrons populate e\textsubscript{π}\textsuperscript{π} (d\textsubscript{xy},d\textsubscript{yz}) creates the possibility of a high-spin (\( S = 3/2 \)) d\textsuperscript{3} ground state with an M–oxo π-bond order of 1. High-spin complexes with (b\textsubscript{2})(e\textsubscript{π}\textsuperscript{π})\textsuperscript{2} configurations must have smaller values of \( \Delta E_{p} \) than those with low-spin d\textsuperscript{3} [(b\textsubscript{2})(e\textsubscript{π}\textsuperscript{π})\textsuperscript{2}] ground states, owing to the presence of an extra M–oxo π-antibonding electron. Even in complexes where the high-spin and low-spin states are degenerate, the high-spin complex will have longer M–oxo bonds and smaller \( \Delta E_{p} \) than the low-spin complex. A forbidden zone of \( \Delta E_{p} \) values emerges from this coupling between electronic configuration and M–oxo bond length. This ambiguity is not present in d\textsuperscript{4} configurations unless population of higher lying σ\textsuperscript{π} orbitals becomes feasible.

Nature knew about the oxo wall when she picked metals for the generation and reduction of oxygen. In their reaction cycles, manganese (PS II) and iron (cytochrome c oxidase and P450) form multiply bonded metal–oxo with dramatically different properties (Figure 3). As we would predict, metal–oxo electronic structure is key: the triply bonded Mn\textsuperscript{V}–oxo is the conjugate base of a superacid, with a pK\textsubscript{a} less than −10. Such a highly electron-deficient metal–oxo could attract an oxygen donor, which in turn would promote redox-coupled O–O bond formation as proposed in one (but not the only) candidate mechanism for oxygen generation from water.

The Fe\textsuperscript{IV}–oxo is much more basic than a Mn\textsuperscript{V}–oxo, owing to the presence of two Fe–oxo π\textsuperscript{π} electrons. Ferryl species of these type are found in compounds I and II of the heme peroxidases. A key determinant of metal–oxo basicity, in addition to the M–oxo π\textsuperscript{π} electron count, is the nature of the ligand \textit{trans} to the oxo. Strongly donating ligands produce more basic oxos than weak donors. The proximal imidazole ligand in peroxidases is an intermediate strength ligand, and there has long been a question of whether compound II in peroxidases was PFe\textsuperscript{IV}–oxo (P = porphyrin) or PFe\textsuperscript{IV}–OH. X-ray and neutron crystallographic studies on ascorbate peroxidase compound II strongly indicate that the preferred configuration is PFe\textsuperscript{IV}–OH. Peroxidase compounds I, however, are best described as P\textsuperscript{+}Fe\textsuperscript{IV}–oxo. The fact that peroxidase compounds II are likely PFe\textsuperscript{IV}–OH species explains their relatively rapid formation from P\textsuperscript{+}Fe\textsuperscript{III}–(H\textsubscript{2}O\textsubscript{2}) precursors. Formation of M–oxo species by oxidation of M–(OH\textsubscript{2}) is extremely sluggish, presumably owing to the necessity to transfer two protons as well as one electron. The
transfer of a single proton and electron to form PFεIV−OH makes for a more facile transformation but does not preclude the intermediacy of P⁺FeIII−(OH).13−17

CYTOCHROME P450

Two π* electrons and an axial cysteine thiolate combine to make the doubly bonded FeIV−oxo of P450 compound II very basic, with a pKₐ of about 12.18,19 It is now widely recognized that the generation of such a basic FeIV−oxo is an essential step in the catalytic cycle of P450. Indeed, Green pointed out that compound I of P450 could extract an electron from a C−H bond only if coupled to proton transfer to the newly formed FeIV−oxo of compound II.19 The high reduction potential of compound I and the basicity of compound II together provide the driving force for the reaction (Figure 4).20

The idealized stoichiometry of cytochrome P450 catalysis predicts that one molecule each of NAD(P)H and O₂ are consumed for each molecule of oxidized substrate produced. Indeed, the prototypical prokaryotic enzyme from Pseudomonas putida (P450cam, CYP101) catalyzes hydroxylation of camphor at the S-exo position in 94% yield relative to NADH consumed.22 Replace camphor with dehydrocamphor, however, and the yield of S-exo-epoxide drops to 75% (relative to NADH). Substrate deuteration also lowers the product yield in CYP101, with the excess electrons producing H₂O from O₂.23,24 This loss of coupling of between NADH/O₂ consumption and product formation has been observed in many P450 enzymes, even with “natural” substrates. The human liver enzyme CYP3A4, found on the endoplasmic reticulum membrane and responsible for a broad range of substrate hydroxylation reactions, including the metabolism of roughly half of therapeutic drugs,25 couples NADH and O₂ consumption to substrate oxidation with barely 10% efficiency.26 Sligar and co-workers identified three branching points in the canonical P450 mechanism that could lead to uncoupling.26,27 The first branch point occurs at the FeIII(O₂−) intermediate, involving competition between delivery of a second electron and O₂− loss (autodissociation). The second branch point occurs at the FeIII(O₂H) intermediate: O−O bond scission leads to ferryl compound I in competition with loss of H₂O₂ (peroxide shunt). The final branch point reflects the competition between substrate oxidation by compound I and two electron transfers to compound I to produce water (oxidase pathway).

We have suggested that reductants internal to the enzyme could protect the enzyme from damage by compound I when reaction with substrate is not possible.28 Intraprotein electron transfer (ET) to ferryl compound I will be a first-order kinetics process with a time constant that is independent of the concentrations of external reductants. The survival time of compound I will depend only on the placement and reduction potential of this antioxidant residue. Redox chains comprised of tyrosine and tryptophan residues (Tyr/Trp chains) could guide oxidizing equivalents (i.e., holes) away from the critical active site, steering them to the enzyme surface where they could be scavenged by soluble reductants (e.g., glutathione, ascorbate, ferrocyanochrome b₅).

Tyr/Trp Chains in P450

We have examined over 90 000 X-ray crystal structures in the RCSB Protein Data Bank (PDB) to identify Tyr/Trp chains in proteins.28 We found that long chains (≥3 residues) occur with highest frequency in the glycosylases and oxidoreductases. The survival time of P450 compound I will depend on the proximity of the nearest Tyr or Trp residue. We determined the shortest heme−Tyr/Trp distances in 134 cytochrome P450 structures with less than 90% sequence identity available in the PDB in April 2018 (Figure 6). The values range from 3.4 to 17.2 Å with a mean of 8.4 Å and standard deviation of 2.9 Å. Although the results should be interpreted cautiously, owing to distortions, owing to the limited sample size, some interesting trends appear in these data. Tyrosine is the redox-active residue closest to the heme in 81% of the bacteria and archaea, whereas in the eukaryotes 71% are tryptophans. Green’s analysis of the high pKₐ of FeIV(OH) in P450 points to the fact that the Gibbs free-energy change for nonproductive Tyr oxidation by compound I is comparable to that for productive activation of an aliphatic C−H bond.19 The energetics are
evenly split between the two pathways, and the net reaction flux will be determined by their relative rates. Productive C−H activation requires positioning of a suitable substrate with the target H atom close to the compound I FeIV−oxo.20 We suggest that the nonproductive antioxidant pathway involves long-range electron transfer from Tyr or Trp. Neglecting small variations in reaction energetics, the key determinant of the internal antioxidant time constant will be the distance from the heme to the nearest Tyr/Trp residue.29−31 If the ferryl fails to oxidize substrate within the prescribed time limit, reducing equivalents from Tyr/Trp residues will rescue the enzyme by regenerating the ferric resting state. The estimated compound I survival time for a Tyr or Trp residue 8.4 Å from the heme (ΔG° = 0.2 eV, λ = 0.8 eV) is about 10 μs; across the full range of heme−Tyr/Trp distances, estimated survival times are as short as 30 ns (3.4 Å) and as long as 140 ms (17.2 Å). Regeneration of the ferric enzyme requires the delivery of a second electron and proton to compound II. The reduction potential of compound II is likely to be near that of compound I,32 so Tyr or Trp residues could again provide the reducing equivalents, although with more sluggish kinetics.13,14

**Protection Pathways**

Our analysis of heme−Tyr/Trp distances (Figure 6) found that values of 7−8 Å are the most common. A large fraction of the 41 structurally characterized eukaryotic P450s are mammalian enzymes that have a Trp residue hydrogen bonded to one of the heme-propionates. This position is occupied by a

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**Figure 5.** Cytochrome P450 catalytic cycle begins with substrate (RH) binding and water displacement in the distal pocket of the ferriheme. Sequenced delivery of electrons, dioxygen, and protons leads to an active intermediate (compound I) responsible for substrate hydroxylation. Three short circuits (dashed arrows) lead to nonproductive dioxygen consumption via autoxidation (A), peroxide shunt (P), and oxidase (O) pathways.

**Figure 6.** Structural database of the RCSB Protein Data Bank contains X-ray crystal structures of 134 cytochromes P450 with sequence identity less than 90%. The histogram illustrates the distribution of the shortest distances between the heme (Fe or members of the porphyrin π-system) and side chain π-system atoms of the redox-active Tyr or Trp residues within this set of P450 structures. The colors indicate the contributions from the three domains: archaea (green), bacteria (blue), eukaryotes (red).
His residue in many of the prokaryotic enzymes. The corresponding residue in human CYP3A4 is Trp126, located 7.2 Å from the heme, and 4.7 Å from surface exposed Tyr99 (PDB ID 1TQN, Figure 7).33 We have analyzed a protection pathway involving hole transfer from the heme via Trp126 and terminating at Tyr99. A hopping map predicts time constants for migration of the hole from compound I to Tyr99 as functions of the driving forces for the two reactions.35 Taking values of $\Delta G^{\ddagger}_1 = -0.1$ eV and $\Delta G^{\ddagger}_2 = 0.1$ eV, we estimate a CYP3A4 compound I survival time of 200 ns. This estimate could be in error by an order of magnitude or more, but it illustrates the tight tolerance for productive substrate oxidation. Once the hole reaches Tyr99, it can be scavenged by the NADH/cytochrome $b_5$ redox system or by soluble reductants such as glutathione.

A primary function of CYP3A4 is xenobiotic metabolism, and its active site binds a broad range of organic substrates. Poor coupling is expected from substrate diversity, and antioxidant rescue pathways are essential for enzyme survival. In this regard, it is noteworthy that cytochrome $b_5$ has been reported to exhibit protective effects for CYP3A4 expressed in E. coli cells.35 Chemical cross-linking studies with human CYP3A4 and cytochrome $b_5$ indicate that P450 residues Lys96 and Lys126 interact with Glu56 on cytochrome $b_5$.36 The two P450 lysine residues span the location of Tyr99 on the CYP3A4 surface (Figure 7), suggesting a route for electron transport along a nonproductive oxidase pathway.

Another human P450 enzyme, CYP11A1, found on the inner mitochondrial membrane, catalyzes the first step in sterol biosynthesis, the conversion of cholesterol to pregnenolone in three turnovers, requiring one O$_2$ and one NAD(P)H molecule per step. This coupling between NAD(P)H consumption and pregnenolone formation is high (>$90\%$),37 and the enzyme is unreactive toward xenobiotics.38 As in CYP102A1 and CYP3A4, one of the heme propionates in CYP11A1 is hydrogen bonded to a tryptophan residue — Trp108, located 7.6 Å from the closest heme atom (PDB ID 3NY9).39 In contrast to the situation for Trp126 in CYP3A4, no redox-active residues are within 10 Å of Trp108. A second tryptophan residue, Trp87, lies above the distal face of the heme at a distance 8.1 Å. Trp87 is the first member of a five-residue chain (Trp87, Trp231, Tyr90, Tyr94, Tyr93) that extends to the enzyme surface with a 3.7 Å average separation distance between residues (Figure 8). Kinetics modeling of this hole transfer pathway suggests that the compound I survival time is about 0.5 μs. Beratan and co-workers examined the electronic coupling strengths between residue pairs in this pathway, reporting values in the range 10–500 cm$^{-1}$.40 They calculated a 1 μs mean first passage time for this pathway.

**HOLE HOPPING IN CYP102A1 (P450BM3) AND CYP119**

The P450 from *Bacillus megaterium* (P450-BM3, CYP102A1) differs from most prokaryotic enzymes in that tryptophan (Trp96) is the closest redox-active residue to the heme. The CYP102A1 X-ray crystal structure reveals that Trp96 is H-bonded to the heme propionate at a distance of 7.2 Å (PDB ID 2J2) in a position closely analogous to that found in many eukaryotic enzymes (Figure 9).41 This enzyme has been the focus of our efforts to study hole hopping reactions in P450s.42,43 We attached a Ru(diimine)$_2^{2+}$ residue to a mutant Cys97 (Ru$_{Cys97}$(CYP102A1)W96) and demonstrated that flash-quench generated Ru$_{Cys97}$(CYP102A1)W96 will oxidize P-(Cys$-S$)$^{3+}$Fe$_{III}$(OH)$_2$ to P(Cys$-S$)Fe$_{IV}$(OH) with a time constant of about 300 μs.43 The kinetics of this reaction indicate that the initial reaction product, (P(Cys$-S$)$^{3+}$)$^{2+}$Fe$_{III}$(OH)$_2$), relaxes by internal proton and electron transfer to yield P(Cys$-S$)-Fe$_{IV}$(OH)$_2$.44

The archaeal P450 enzyme from *Sulfolobus acidocaldarius* (CYP119, PDB ID 1HO7)45,46 has a His residue (His76) at the same location as Trp96 in CYP102A1. We prepared a conjugate with Ru(diimine)$_2^{2+}$ bound to a mutant Cys77 residue in CYP119 and found that flash-quench generated Ru$_{His76}$(CYP119)H76 does not oxidize...
sulfur atom is commonly, although not exclusively, found. This coordination environment produces the intense blue color of the CuII forms, and leads to CuII formal potentials ranging from 184 mV vs NHE for stellacyanin to 680 mV for rusticyanin. Of 30 structurally characterized cupredoxins, none has Tyr or Trp residues within 5 Å of the copper center. The only cupredoxin with a Tyr or Trp residue near the active site is the iso-2 azurin from *Methylomonas* sp. (strain J) (PDB ID 1CUO).49 Tyr114 is 5.03 Å from Cu.

Multicopper oxidase enzymes (MCO) are constructed from 2 or 3 cupredoxin domains, one of which retains a type 1 copper. A trinuclear Cu (TNC) active site is at the interface between two of the domains, and it is here that oxygen is reduced to water. A survey of 25 X-ray crystal structures of 3-domain MCOs reveals that a Trp or Tyr/Trp pair is found adjacent to the TNC in all but two of these enzymes. In the *Thermus thermophilus* enzyme, Trp133 is just 3.6 Å from one of the Cu centers in the TNC (PDB ID 2YAE, Figure 10).52 This residue could play a protective role for the enzyme, analogous to that postulated for the cytochromes P450. Under normal circumstances, those reducing equivalents could be delivered by four CuI centers. In cases where four electrons are not available, Trp133 could supply an electron to prevent formation of damaging reactive oxygen species.

**Figure 10.** Trp133 is positioned 3.6 Å from a TNC Cu center in the MCO from *Thermus thermophilus* (PDB ID 2YAE). Reduction of oxygen to water requires four electrons. Under normal circumstances, those reducing equivalents could be delivered by four CuI centers. In cases where four electrons are not available, Trp133 could supply an electron to prevent formation of damaging reactive oxygen species.

**Figure 9.** Illustration of the structural similarity of Ru-modified CYP102A1 (PDB ID 3NPL) and CYP3A4 (green, PDB ID 1TQN). Trp96 in CYP102A1 mediates high potential hole transfer from the heme to photochemically generated Ru(diimine).35. Trp126 in CYP3A4 occupies the same site as Trp96 in CYP102A1; Tyr99 in CYP3A4 is adjacent to the Ru binding site in CYP102A1.
FIGURES

Figure 1. A schematic diagram of the relationship between the redox potential and the energy of the O2 molecule.

Figure 2. A detailed view of the active site of the enzyme, showing the key residues involved in the O2 activation process.

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