Nature of the Ferryl Heme in Compounds I and II

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Heme enzymes are ubiquitous in biology and catalyze a vast array of biological redox processes. The formation of high valent ferryl intermediates of the heme iron (known as Compounds I and Compound II) is implicated for a number of catalytic heme enzymes, but these species are formed only transiently and thus have proved somewhat elusive. In consequence, there has been conflicting evidence as to the nature of these ferryl intermediates in a number of different heme enzymes, in particular the precise nature of the bond between the heme iron and the bound oxygen atom. In this work, we present high resolution crystal structures of both Compound I and Compound II intermediates in two different heme peroxidase enzymes, cytochrome c peroxidase and ascorbate peroxidase, allowing direct and accurate comparison of the bonding interactions in the different intermediates. A consistent picture emerges across all structures, showing lengthening of the ferryl oxygen bond (and presumed protonation) on reduction of Compound I to Compound II. These data clarify long standing inconsistencies on the nature of the ferryl heme species in these intermediates.

Aerobic organisms have evolved to utilize the intrinsic oxidizing power of dioxygen. In a large majority of cases, this so-called “activation” of oxygen is catalyzed by a metal center (usually iron or copper) buried within a protein structure (1). The large and biologically diverse family of catalytic iron-containing heme proteins is one such group of enzymes capable of oxygen activation. Oxygen activation by heme is achieved through formation of highly oxidized iron intermediates, which are known as Compound I and Compound II. Compound I is formally oxidized by two (electron) equivalents above the ferric heme resting state; Compound II is the reduced form of Compound I and thus oxidized by only one equivalent. Nature uses these intermediates for a large number of quite different, and sometimes difficult, biological oxidations. The most well cited examples are in the cytochrome P450s, nitric oxide synthases, and heme peroxidases (2–5). It appears that there are two possible routes for formation of Compound I, a direct reaction with hydrogen peroxide (e.g. the peroxidase enzymes) or reaction with dioxygen followed by a further reduction of the heme by a suitable reductase (e.g. the P450s, NO synthase (Scheme 1)). However, the shared heme structure used in all of these enzymes, together with the similarity of their reactions with dioxygen or dioxygen derivatives, make it highly likely that these intermediates and their mechanisms of formation are a defining feature across the family.

It is no surprise, therefore, that elucidation of the structure of these heme intermediates has provided an important focus and is the source of continuing high profile comment and debate (see for example Ref. 6). However, the quest to unravel the structure and properties of Compounds I and II has not been at all straightforward, not least because of the experimental difficulties associated with observing such metastable species. A key challenge has been to clarify the precise nature of the ferryl heme species as the literature reports conflicting bond lengths measured for the Compound I or Compound II intermediates in different proteins. The main experimental approaches have used EXAFS3 data and crystallography, but an inconsistent picture emerges (see Ref. 7 for a recent summary). This has made it difficult to determine the bonding interactions with confidence (Fe(IV) = O versus Fe(IV)–OH, the latter being a longer bond). This is made yet more complicated by the fact that many previous crystallographic studies are likely to have been affected by photoelectron reduction in the x-ray beam so that deduced structures are unlikely to represent “pure” species. Although the effect of x-ray photoreduction on the reaction intermediates has been explored in detail for horseradish peroxidase (HRP) (8), a precise structural characterization of these intermediates has not been achieved in any other enzymes.

In this work, we present crystal structures of both the Compound I and the Compound II intermediates in two different peroxidase enzymes, cytochrome c peroxidase (CcP) and the closely related ascorbate peroxidase (APX). The structures provide reliable measurements of bond distances and in both cases indicate a lengthening of the bond on reduction that is presumed to be coupled to protonation. The likely mechanisms of proton delivery to the transient ferryl heme are also revealed. The information is relevant to our understanding of
oxygen activation across the family of heme enzymes and is discussed in this more general context.

**EXPERIMENTAL PROCEDURES**

**Protein Purification and Expression**—The MKT variant of CcP (9) was prepared as previously described but cloned into the expression plasmid pLEICS-03 carrying kanamycin resistance and a tobacco etch virus-cleavable N-terminal His tag sequence and expressed in BL21 DE3 Gold (Fisher Scientific). Expression and purification of APX were carried out as previously described (10, 11).

Both CcP and APX were crystallized as previously described (12, 13). CcP crystals were grown in 50 mM potassium phosphate, pH 6.0, containing (±)-2-methyl-2,4-pentanediol (30% v/v), and APX crystals were prepared from Li2SO4 (2.25 M) containing HEPES (0.1 M), pH 8.3. Compound I of CcP was formed by soaking crystals of ferric enzyme in freshly prepared H2O2 (20 mM) for 5 min, and crystals were then flash-frozen at 100 K. Compounds II and III of APX were prepared by soaking the ferric enzyme in H2O2 (20 mM) for 5 min or H2O2 (20 mM) for 5 min, respectively, and frozen as above. Crystals were stored in liquid nitrogen.

**Monitoring of Photoreduction**—Changes in UV-visible absorbance spectra during x-ray exposure were monitored at the European Synchrotron Radiation Facility (ESRF) ID14-2 using an on-line spectrophotometer (OCEAN OPTICS DH 2000 light source and HR 2000 detector). UV-visible spectra of exposed crystals used to solve the structure of CcP Compound II, APX Compound I, ferrous CcP, and APX were measured using a 4DX single crystal microspectrophotometer with a Shamrock SR-163 spectrograph and Newton CCD camera (Andor Technology).

**Crystallographic Data Collection and Analysis**—For the structure of CcP intermediates, x-ray diffraction data were collected in-house at 100 K using CuKα radiation (λ = 1.5418 Å) from a Rigaku RU2HB x-ray generator with a copper anode and Xenocs multilayer optics and measured with an R-AXIS IV detector. For APX, diffraction data were collected on beam line I04 (wavelength λ = 0.6 Å) at Diamond Light Source, Harwell, UK using an ADSC Q315 CCD detector, at 100 K. CcP data were indexed, integrated, merged, and scaled, respectively, using MOSFLM, SORTMTZ, and SCALA as part of the CCP4 suite (14, 15); APX data were integrated, merged, and scaled using XDS (16). In all cases, 5% of the data were flagged for the calculation of R_free and excluded from subsequent refinement. Data collection statistics are shown in Table 1.

**Multicrystal Data Analysis**—The CcP Compound I structure was solved by merging the first 9° of data from 10 different crystals. This corresponds to a maximum absorbed dose of 0.020 MGY (calculated with RADDose (17)). We established the amount of data that could be collected from each crystal before the effects of photoreduction showed empirically by truncating the data after various doses. We determined that no detectable change in the observed structure was seen with this (or a lesser) dose but was apparent after a higher dose. The APX Compound II and Compound III structures were solved from three crystals as it was found that by using 0.6 Å radiation, the photoelectron effect is significantly reduced (18), allowing at least 15° to be collected before there was any indication of photoreduction. This corresponds to an absorbed dose of 0.028 MGY (calculated with RADDose (17)). Compound I of APX was produced by photoreduction of Compound III (absorbed dose ~0.15 MGY) and verified by UV-visible single crystal spectrophotometry. Diffraction data for Compound I were also obtained from three crystals (using the first 15° after the photoreduction). Compound II of CcP was obtained by the photoreduction of Compound I (absorbed dose of ~0.15 MGY) and verified by UV-visible single crystal spectrophotometry. Diffraction data were collected following photoreduction by merging together the first 9° of data from 10 crystals.

**Refinement**—Crystallographic refinement initially used REFMAC5 (19) from the CCP4 suite (15). The structure of Compound I of CcP was refined from the 1.70 Å wild-type CcP structure (20) (1ZBY), whereas the structure of Compound II of APX was refined from the 1.45 Å ferric APX structure (13) (1OAG). To ensure the unbiased determination of the iron-oxygen distances, the entire protein structure was refined with the ferryl oxygen atom omitted, and the ferryl oxygen was first fitted to the peak of the $F_o - F_c$ difference.
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map and refined in real space using Coot (21). The complete models were then refined with SHELX (22), allowing the estimation of individual atomic positional uncertainties (estimated standard uncertainty (ESU)).

RESULTS

We have obtained structures of five reaction intermediates, Compounds I and II for bothCcP and APX and Compound III (ferrous-oxy) of APX. Compound I of CcP and Compounds II and III of APX are isolatable species, and thus we are able to obtain crystals of these directly by reaction with \( \text{H}_2\text{O}_2 \). The structure of CcP Compound II was obtained indirectly, by photoreduction of its Compound I; likewise APX Compound I was also obtained indirectly, through photoreduction of Compound III.

CcP Compound I—The structure of the Compound I intermediate of CcP has been determined using a multicrystal approach (using only a small percentage of the total data set (8)) to allow the collection of diffraction data before it is affected by photoreduction (Fig. 1A). Data and refinement statistics are shown in Table I. The authenticity of this Compound I species was confirmed using single crystal microspectrophotometry (Fig. 2A), which shows the typical Compound I peaks (530, 560, and 632 nm) in the visible region and compares them with previously published spectra (530, 560, and 630 nm (23)) and with our spectra in solution under the same conditions (i.e. using the same ratio of [enzyme]:[\( \text{H}_2\text{O}_2 \)] (Fig. 2A)). The structure (Fig. 1A) reveals an electron density peak for the oxygen atom at 1.63 Å from the iron. This is considerably shorter than previous crystallographic estimates of the Compound I bond length in CcP (1.87 Å (24) and 1.7–2.0 Å (25)). The ESU of the iron and oxygen positions calculated by full matrix inversion (22) are 0.017 and 0.066 Å, respectively. When compared with Compound I, this clearly indicates a lengthening of the Fe–O ferryl bond in the Compound II structure. The similarity of the single crystal and solution spectra for Compound II (Fig. 2B) and the fact that the bond length for the Compound II species is much shorter than that for either the published ferric APX (2.08 Å) or the ferrous APX (2.20 Å below) give confidence that the assignment of the structure as a Compound II species is correct.

The ferryl iron also moves out of the heme plane by 0.15 Å toward the distal histidine (when compared with the ferric enzyme), and the proximal histidine (His-163) also shifts in the same direction, in the same way as was observed for Compound I of CcP above. The side chain of the distal arginine is seen in both the in (as in Compound I) and the out (as in the ferric enzyme) conformations.

As for CcP Compound I above, photoreduction of Compound II of APX is observed during exposure to the x-ray beam, leading to the formation of ferrous APX (shown by peaks emerging at 554 nm and a decrease at 534 nm (Fig. 2B)) for which we have also obtained a structure (Fig. 3B). The distance between the Fe(II) and the oxygen of water is now 2.2 Å, which is much longer than those for any of the Compound I or Compound II species above.

We sought additional structural information on the Compound I and Compound II bond lengths observed above in the corresponding Compound I and Compound II derivatives of APX and CcP, respectively. This is described below.

APX Compound I—The Compound I intermediate of APX converts rapidly to Compound II (28), and we are thus unable to isolate this form in the crystal by soaking with peroxide. We can, however, access the Compound I intermediate through the ferrous-oxy (Fe(II)–O\(_2\)) species (Compound III), which is used in other catalytic heme enzymes (such as the P450s) as a route to formation of Compound I (Scheme 1). In peroxydases, the ferrous-oxy intermediate can be formed from reaction of Compound II with excess peroxide (8); photoreduction of the ferrous-oxy (Compound III) species thus formed in the crystal converts it to the desired Compound I. The ferrous-oxy species is analogous to the ferric-hydroper-
oxide species (sometimes referred to as Compound 0, Fe(III)–OOH) that precedes Compound I and can therefore also report on this transient precursor.

The structure of this ferrous-oxy (Compound III) species is shown in Fig. 1C. An electron density peak larger than those observed for the Compound I and Compound II structures is...
TABLE 1

<table>
<thead>
<tr>
<th>Data collection and refinement statistics</th>
<th>CpP</th>
<th>Compound I</th>
<th>Compound II</th>
<th>Ferrous</th>
<th>APX</th>
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<tr>
<td>Data collection&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
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<td>P2&lt;sub&gt;1&lt;/sub&gt;,2,2</td>
<td>P2&lt;sub&gt;1&lt;/sub&gt;,2,2</td>
<td>P2&lt;sub&gt;1&lt;/sub&gt;,2,2</td>
<td>P4&lt;sub&gt;2&lt;/sub&gt;,2</td>
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<td>Cell dimension a, b, c (Å)</td>
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<td>51.05, 75.19, 106.95</td>
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<td>Resolution (Å)</td>
<td>1.67 (1.76–1.67)</td>
<td>1.67 (1.76–1.67)</td>
<td>1.69 (1.78–1.69)</td>
<td>1.55 (1.59–1.55)</td>
<td>1.50 (1.54–1.50)</td>
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<td>R&lt;sub&gt;work&lt;/sub&gt;/R&lt;sub&gt;free&lt;/sub&gt; (%)</td>
<td>10.9 (3.8)</td>
<td>5.4 (2.4)</td>
<td>15.8 (4.0)</td>
<td>13.1 (2.0)</td>
<td>14.6 (4.4)</td>
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<td>Multiplicity</td>
<td>92 (81)</td>
<td>83.5 (75.8)</td>
<td>99.4 (97.2)</td>
<td>87.9 (97.2)</td>
<td>88.6 (84.7)</td>
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<td>Completeness (%)</td>
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<td>2.1 (2.0)</td>
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<td>3.7 (1.4)</td>
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<td>6.8 (60.0)</td>
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<td>Resolution (Å)</td>
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<td>33.15–1.69</td>
<td>57.96–1.55</td>
<td>27.7–1.68</td>
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<td>44336</td>
<td>50009</td>
<td>37890</td>
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<td>R&lt;sub&gt;work&lt;/sub&gt;/R&lt;sub&gt;free&lt;/sub&gt; (%)</td>
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<td>0.052</td>
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<td>0.044</td>
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<td>Bond lengths (Å)</td>
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<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Bond angles (Å)</td>
<td>0.024</td>
<td>0.025</td>
<td>0.047</td>
<td>0.026</td>
<td>0.024</td>
</tr>
<tr>
<td>r.m.s.</td>
<td></td>
<td></td>
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<sup>a</sup> Data collected at home source at wavelength 1.5418 Å merging together the first 9° of data from 10 crystals.
<sup>b</sup> Data were collected at Diamond Light Source beam line IO4 at wavelength 1.0 Å merging the first 15° of data from three crystals. As the atomic cross-section decreases with the wavelength (8), the use of a high energy beam considerably reduced the photoelectric effect and thus decreased the number of crystals needed.
<sup>c</sup> Outer bin data in brackets.
<sup>d</sup> Calculated by the maximum likelihood method (19).
<sup>e</sup> r.m.s., root mean square.

A–D, comparisons for CpP (A and D) and for APX (B and C). A, the single crystal spectrum of CpP Compound I collected before x-ray exposure (solid line, maxima: 530, 560, and 632 nm) and after collection of a full dataset (dose > 0.35 Mgy, dash-dotted line, maxima: 542, 559, and 585th nm corresponding to reduced heme) when compared with the spectrum of CpP Compound I obtained in solution under the same conditions (dotted line, maxima: 530, 560, and 630th nm). Abs, absorbance. B, the spectrum of APX Compound I collected before x-ray exposure (solid line, maxima: 531, 558 nm) and after exposure to a dose of 0.3 Mgy (dash-dotted line, maxima: 537<sup>th</sup>, 554, and 580<sup>th</sup> nm corresponding to reduced APX (compare 555 and 583 nm (29))) when compared with the solution spectrum of APX Compound II obtained under the same conditions (dotted line, maxima: 530, 559 nm (compare 529 and 560 nm (27))). C, spectra of APX Compound III (dash-dotted line, maxima: 540 and 575 nm) and APX Compound I (solid line, maxima 535, 571<sup>th</sup> and 623 nm) obtained by photoreduction of APX Compound III after exposure to a dose of 0.15–0.2 Mgy. Appearance of the shoulder at 585 nm and slight red shift of the β band from 530 to 533 nm indicate the presence minor amounts of ferrous heme formed during photoreduction.

FIGURE 2. Comparison of UV-visible spectra in solution and from single crystal microspectrophotometry. A–D, comparisons for CpP (A and D) and for APX (B and C). A, the single crystal spectrum of CpP Compound I collected before x-ray exposure (solid line, maxima: 530, 560, and 632 nm) and after collection of a full dataset (dose > 0.35 Mgy, dash-dotted line, maxima: 542, 559, and 585<sup>th</sup> nm corresponding to reduced heme) when compared with the spectrum of CpP Compound I obtained in solution under the same conditions (dotted line, maxima: 530, 560, and 630<sup>th</sup> nm). Abs, absorbance. B, the spectrum of APX Compound I collected before x-ray exposure (solid line, maxima: 531, 558 nm) and after exposure to a dose of 0.3 Mgy (dash-dotted line, maxima: 537<sup>th</sup>, 554, and 580<sup>th</sup> nm corresponding to reduced APX (compare 555 and 583 nm (29))) when compared with the solution spectrum of APX Compound II obtained under the same conditions (dotted line, maxima: 530, 559 nm (compare 529 and 560 nm (27))). C, spectra of APX Compound III (dash-dotted line, maxima: 540 and 575 nm) and APX Compound I (solid line, maxima 535, 571<sup>th</sup> and 623 nm) obtained by photoreduction of APX Compound III after exposure to a dose of 0.15–0.2 Mgy. Appearance of the shoulder at 585 nm and slight red shift of the β band from 530 to 533 nm indicate the presence minor amounts of ferrous heme formed during photoreduction.

clearly seen and interpreted as a dioxygen species. This is confirmed by single crystal spectrophotometry (peaks at 548 and 575 nm (Fig. 2C)). There are hydrogen-bonding interactions from the O<sup>2</sup> of the bound ligand to N<sup>ε</sup> of His-42 (2.92 Å), N<sup>ε</sup> of Trp-41 (2.70 Å), and N<sup>ε</sup> of Arg-38 (2.65 Å). In this case, Arg-38 (equivalent to Arg-48 in CpP) is observed occupying both the in and the out positions, and there is a water molecule, seen adjacent to Trp-41 (W1 in Fig. 1C), which shifts away from the heme to accommodate the bound O<sub>2</sub> species. Photoreduction leads to cleavage of the O–O bond and the ultimate formation of Compound I (8). We observe that after a dose of ~0.15 Mgy, a structure is seen in which only a single atom of oxygen is bound above the iron. Single crystal spectrophotometry (Fig. 2C) confirms that this is predominately Compound I (27), and the structure is shown in Fig. 1D. In this structure, the iron-oxygen bond length is 1.73 Å, which is slightly longer than that obtained for the Compound I structure of CpP above (most likely because the structure still contains some O<sub>2</sub> bound to the heme (residual ferrous-oxy-heme)) but still shorter than previous crystallographic measurements from CpP (Table 2). The ESUs of the iron and oxygen positions (22) are 0.016 and 0.09 Å, respectively.

CpP Compound II—As for APX Compound I, Compound II of CpP is not stable enough to be trapped and isolated directly in the crystal but can be accessed indirectly through photoreduction of Compound I. The spectrum of CpP Compound II is very similar to that of Compound I (the only difference between the two species being that the Trp-191 radical has been reduced in Compound II (23)), which makes unambiguous identification of Compound II in the crystal more difficult, but the intensity of the α- and β-bands decreases on reduction of Compound I to Compound II (23). To obtain a structure for CpP Compound II, we have used diffraction data after an exposure of 0.15–0.20 Mgy where the α- and β-bands have decreased, but formation of ferrous heme is still negligi-
ble (as monitored by the presence of the characteristic 585 nm peak for the ferrous species (Fig. 2A)). Fig. 1E shows the structure of the Compound II species thus obtained; single crystal spectra (Fig. 2D) confirm the assignment and are in agreement with previous spectra for Compound II (23).

In this case, the Fe–O distance is observed to clearly increase and is now longer (1.83 Å) than that for both CcP Compound I (1.63 Å) and APX Compound I (1.73 Å) above and almost identical to that seen in APX Compound II (1.84 Å). The ESUs of the iron and oxygen atom positions calculated by full matrix inversion are 0.019 and 0.159 Å, respectively. Arg-48 mostly remains in the in position, although some positive electron density is observed that is consistent with a low occupancy in the out position.

**DISCUSSION**

The nature of the Compound I and Compound II intermediates in different heme enzymes has been the subject of intense and sometimes confusing debate (see Ref. 30 for a recent discussion). X-ray crystallographic studies varied in their conclusions on bond length probably because these early structures were photoreduced. A summary of bond lengths from various studies is presented in Table 2.

Here we present the structures of Compound I of CcP and Compound II of APX, neither of which is affected by photoreduction. We also present the corresponding (and consistent) structures of Compound I of APX and Compound II of CcP. All of the structures obtained here are summarized and compared in Scheme 2. The two proteins share high sequence identity, and their overall structures are very similar (13, 31). This allows a direct and meaningful comparison of the structures of the intermediates in two different proteins. Together with the structures of the ferrous and ferrous-oxy (the latter for APX only) species, this provides a detailed comparative picture of the key redox states (Scheme 2).

**TABLE 2**

<table>
<thead>
<tr>
<th>Protein</th>
<th>Compound I</th>
<th>Compound II</th>
<th>Reference</th>
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<tr>
<td>Horseradish peroxidase</td>
<td>1.6&quot;a</td>
<td>1.6&quot;a</td>
<td>(41)</td>
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<td></td>
<td>1.64&quot;a</td>
<td>1.64&quot;a</td>
<td>(37)</td>
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<td>1.67&quot;a</td>
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<td></td>
<td>1.72&quot;a</td>
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<td>(8)</td>
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<td>Cytochrome c peroxidase</td>
<td>1.67&quot;a</td>
<td>1.83b</td>
<td>(35)</td>
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<tr>
<td></td>
<td>1.87b</td>
<td>2.00c,d</td>
<td>(24)</td>
</tr>
<tr>
<td></td>
<td>1.73b</td>
<td></td>
<td>(32)</td>
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<tr>
<td>Ascorbate peroxidase</td>
<td>1.63b</td>
<td>1.84b</td>
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<tr>
<td>Chloroperoxidase</td>
<td>1.65&quot;a</td>
<td>1.82c</td>
<td>(39, 43)</td>
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<td>Myoglobin</td>
<td>1.69&quot;a</td>
<td>1.92b</td>
<td>(47)</td>
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<td>1.92b</td>
<td>1.92b</td>
<td>(40, 48)</td>
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<tr>
<td>Cytochrome P450 (CYP 119)</td>
<td>1.82c</td>
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<td>(30)</td>
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*a* From EXAFS.

*b* From x-ray crystallography.

*c* There is an early structure for Compound I of CcP (26) but at low resolution (2.5 Å).

*d* In this case, a range of bond lengths is given (1.7–2.0 Å) depending on the refinement.

4 We considered the possibility that the Compound II structures that we present are, in fact, another species such as ferric or ferric-hydroxide, but the microspectrophotometry data are not consistent with such an assignment (ferrous heme is similarly eliminated). The spectrum of ferric APX (29) is easily distinguished from that of Compound II. Our single crystal spectra for CcP Compound II also agree with literature values (23) and with the corresponding spectra that we have measured in solution (Fig. 2) and give confidence of assignment.

In this case, the Fe–O distance is observed to clearly increase and is now longer (1.83 Å) than that for both CcP Compound I (1.63 Å) and APX Compound I (1.73 Å) above and almost identical to that seen in APX Compound II (1.84 Å). The ESUs of the iron and oxygen positions (22) are 0.017 and 0.065 Å, respectively. Arg-48 mostly remains in the in position, although some positive electron density is observed that is consistent with a low occupancy in the out position.
spectra give a high degree of confidence that the structure corresponds to Compound I. Our crystallographic analyses for this ferryl intermediate in Compound I of CcP measures the bond length as 1.63 Å, firmly in the realm of an unprotonated iron–oxo double bond. It is substantially shorter than a previous estimate of 1.87 Å (24), but this structure was affected by photoreduction. This structure can be compared directly with that of APX Compound I. For APX, Compound I is not stable, so its structure necessarily has been obtained indirectly, from photoreduction of the ferrous-oxo intermediate. Nevertheless, the single crystal spectra for the APX Compound I structure are consistent with published data in solution, and the bond lengths in the two Compound I structures are in good agreement; for APX, the bond is slightly longer (1.73 Å) than for CcP (1.63 Å), most likely from the presence of residual ferrous-oxo heme (which would increase the apparent bond length). Both bond lengths for Compounds I of CcP and APX are shorter than other estimates (24, 25). Both of our Compound I structures show an observed bond length consistent with an unprotonated ferryl (Fe(IV)=O) heme species.

It is worth stating here that the Compound I intermediates of APX and CcP are not exactly the same. CcP Compound I contains a tryptophan radical (Trp-191) as the site of the second oxidizing equivalent (33). APX contains the equivalent tryptophan residue (Trp-179) but does not use it; instead, APX Compound I contains a porphyrin π-cation radical, and its spectrum thus differs from that of CcP Compound I. Our data suggest that this difference in the location of the second oxidizing equivalent does not affect the nature of the ferryl heme species.

Nature of the Ferryl Heme in Compound II—As we discuss below, the nature of the ferryl heme in Compound II has been more controversial. We have presented two Compound II structures. Of them, the APX Compound II structure is the most reliable; APX Compound II is well documented by us (27) and others (34) as an isolatable species. The single crystal spectra that we present for APX Compound II unambiguously
confirm this assignment. For Compound II of APX, the bond length is clearly observed as longer (1.84 Å) than for either of the Compound I structures, consistent with a protonated Fe(IV)–OH (single) bond. This is confirmed in the Compound II structure of CcP; in this case, the structure is obtained indirectly (by photoreduction), but the bond also lengthens (to 1.83 Å). Our experiments provide evidence for lengthening of the bond on reduction of Compound I to Compound II in both enzymes, and the data are consistent with protonation of the ferryl heme on reduction.

Implications for Other Heme Ferryl Species—Much, but not all, of the confusion in this area has arisen from crystal structures of ferryl heme species that were reduced by the x-ray beam. For CcP Compound I, the EXAFS and most of the other data now seem to support an unprotonated ferryl-oxo species (32, 35, 36); this is in agreement with our analyses for both CcP Compound I and APX Compound I. Compounds I of HRP (8, 37–39) and chloroperoxidase (39) are also not protonated. There seems to be a consensus, therefore, for the Compound I species across several peroxidases.

The picture for Compound II has been even less clear-cut. Much of the information is available for HRP, for which some data favor an unprotonated ferryl unit (7, 36, 37, 39, 41), whereas other work (8, 38) favors a long bond (protonated oxygen). There is only one published structure (8), the reliability of which has been questioned (36, 42, 43). For chloroperoxidase, EXAFS data support a long bond, and thus protonated oxygen, in Compound II (39, 43). Because of this, it has been suggested (42, 44–46) that thiolate ligation is a necessary requirement for protonation of the ferryl heme unit and is thus a unique feature of thiolate-ligated hemes. This would also include the thiolate-ligated P450s, for which there is evidence of a long bond in Compound II (30) but for which characterization of the ferryl heme species has been especially troublesome. The idea that only thiolate-ligated heme proteins are able to form protonated ferryl species is not consistent with our observations for Compounds II of APX and CcP, both of which contain imidazole as axial ligand. Our data suggest that the bond lengthening observed in chloroperoxidase (39, 43) and HRP (8) on reduction of Compound I to Compound II is also observed in APX and CcP.

Proton Delivery—Any heme enzyme that is using Compound I as an obligate intermediate formally requires two protons for reduction of Compound I back to ferric heme (and concomitant release of the ferryl oxygen atom as water). The source of these protons therefore raises interesting questions. In the case of the peroxidases examined here, the distal arginine appears as a prime suspect. The various structures for both enzymes presented here show different orientations of the distal arginine within the active site (in and out), and in some structures (e.g. Compounds I and II of APX), both orientations are observed. For the in orientation, this arginine side chain is close enough to donate a proton to the ferryl oxygen (Fig. 1). The close hydrogen-bonding interaction clearly supports the idea that this distal arginine side chain can potentially provide a proton to the ferryl oxygen of Compound I on reduction in both CcP and APX. This would be consistent with the observation that the distal arginine is conserved in all peroxidases.

Overall Conclusions—Compound I formation is implicated as a common intermediate in many catalytic heme enzymes but, as we show in Scheme 1, there is more than one mechanism for its formation, and different heme enzymes use different preferred pathways. Compound I in P450s is highly unstable and has been difficult to capture even using fast spectroscopic methods; the corresponding intermediate has been yet more elusive in the NO synthases so that reliable crystal structures for Compound I intermediates in these enzymes have yet to emerge. Our structures, summarized in Scheme 2, show a consistent picture for the ferryl heme species across two different enzymes. In view of the mechanistic similarities that are a hallmark of heme enzyme chemistry, it is very likely that this snapshot of peroxidase structures represents a more general picture across the wider family and is thus likely to be relevant to the development of our understanding of other, less well characterized, catalytic heme enzymes.

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Addendum—While this work was being written up for publication, a related study appeared (32) in which this 1.87 bond length in CcP has been reassessed. The structure reports an Fe–O distance of 1.73 Å for Compound I of CcP, which is longer than the distance (1.63 Å) reported here (the dose received in these experiments was approximately one-third more than the dose reported here) but still consistent with an unprotonated ferryl-oxo species.

REFERENCES
Nature of Ferryl Heme in Compounds I and II

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Nature of the ferryl heme in Compounds I and II.
Andrea Gumiero, Clive L. Metcalfe, Arwen R. Pearson, Emma Lloyd Raven, and Peter C. E. Moody

PAGE 1263:

The legend to Fig. 1 should read as follows.

FIGURE 1. Stereo images of the crystal structures of the ferryl heme intermediates of CcP and APX. A–E, CcP Compound I (A), APX Compound II (B), APX Compound III (ferrous-oxy) (C), APX Compound I (D), and CcP Compound II (E), showing electron density maps calculated with coefficients 2Fo−Fc (contoured at 2σ, shown in blue) and the Fo/Fc map (contoured at 4σ, shown in green) calculated after refinement omitting the oxygen. Oxygen atoms are shown as red spheres, the heme is in red, and the iron is shown as an orange sphere. Key residues are labeled.

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A novel non-SET domain multi-subunit methyltransferase required for sequential nucleosomal histone H3 methylation by the mixed lineage leukemia protein-1 (MLL1) core complex.
Anamika Patel, Valarie E. Vought, Venkatasubramanian Dharmarajan, and Michael S. Cosgrove

PAGE 3363:

In Table 1, a factor of 10^4 was incorrectly inserted in front of the units h^-1 and μM^-1 h^-1 for the apparent Kcat and Kcat/Km values. As a result, the reported values in Table 1 underrepresent the apparent Kcat values by a factor of 10^4. Our error does not affect the conclusions of the article. We apologize for any confusion that this error may have caused. The corrected Table 1 is shown below.

TABLE 1
Summary of apparent kinetic parameters for WRA and WRAD complexes

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>K_m</th>
<th>k_cat</th>
<th>k_cat</th>
<th>K_cat</th>
<th>K_cat/K_m</th>
</tr>
</thead>
<tbody>
<tr>
<td>WRA</td>
<td>13.7 ± 3.7</td>
<td>2382 ± 269</td>
<td>20</td>
<td>70</td>
<td>1.5</td>
</tr>
<tr>
<td>WRAD</td>
<td>7.9 ± 1.7</td>
<td>338 ± 54</td>
<td>18</td>
<td>30</td>
<td>2.3</td>
</tr>
</tbody>
</table>

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