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Characterization of Dehaloperoxidase Compound ES and Its Reactivity with Trihalophenols†

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ABSTRACT: Dehaloperoxidase (DHP), the oxygen transport hemoglobin from the terebellid polychaete Amphitrite ornata, is the first globin identified to possess a biologically relevant peroxidase activity. DHP has been shown to oxidize trihalophenols to dihaloquinones in a dehalogenation reaction that uses hydrogen peroxide as a substrate. Herein, we demonstrate that the first detectable intermediate following the addition of hydrogen peroxide to ferric DHP contains both a ferryl heme and a tyrosyl radical, analogous to Compound ES of cytochrome c peroxidase. Furthermore, we provide a detailed kinetic description for the reaction of preformed DHP Compound ES with the substrate 2,4,6-trichlorophenol and demonstrate the catalytic competency of this intermediate in generating the product 2,4-dichloroquinone. Using rapid-freeze-quench electron paramagnetic resonance spectroscopy, we detected a $g \approx 2.0058$ signal confirming the presence of a protein radical in DHP Compound ES. In the absence of substrate, DHP Compound ES evolves to a new species, Compound RH, which is functionally unique to dehaloperoxidase. We propose that this intermediate plays a protective role against heme bleaching. While unreactive toward further oxidation, Compound RH can be reduced and subsequently bind dioxygen, generating oxyferrous DHP, which may represent the catalytic link between peroxidase and oxygen transport activities in this bifunctional protein.

The terebellid polychaete Amphitrite ornata contains an abundant coelomic hemoglobin (I) that has been named dehaloperoxidase (DHP)† (2). In the presence of hydrogen peroxide, this globin catalyzes the two-electron oxidation of a trihalophenol substrate to yield a dihaloquinone (Scheme 1). Although DHP is dependent upon the pH and substrate employed, kinetic assays demonstrate that DHP is 1–2 orders of magnitude faster than myoglobin (Mb) in its ability to dehalogenate substrate (3), and only 1 order of magnitude slower than horseradish peroxidase (HRP) (4). Thus, DHP

Scheme 1: Reaction of DHP with Trihalogenated Phenolate and Hydrogen Peroxide Yields Quinone Products

is an anomaly: it functions as the oxygen transport protein in A. ornata despite having a low degree of sequence homology with other hemoglobins (5–8) yet exhibits significant peroxidase activity approaching that of HRP even though it possesses neither structural nor sequence homology with any known peroxidase.

Despite a number of studies on DHP, how this bifunctional protein can act as both a hemoglobin and a peroxidase is still not understood (1, 2, 4). Peroxidases generally function via the Poulos–Kraut mechanism (9) in which $\text{H}_2\text{O}_2$ reacts with a ferric heme to form Compound I, the iron(IV)–oxo porphyrin $\pi$-cation radical species that is formally oxidized by two electrons relative to the ferric resting state. The majority of peroxidases regenerate the resting form of the enzyme via two sequential one-electron substrate oxidations ($2\text{AH} + \text{H}_2\text{O}_2 \rightarrow 2\text{A}^+- + 2\text{H}_2\text{O}$), proceeding through Compound II, an iron(IV)–oxo species that is one-electron-oxidized when compared to the ferric state. Furthermore, DHP (and HRP) has been shown to perform the net two-electron oxidation of phenols ($X=\text{I, Br, Cl, F}$) is an anomaly: it functions as the oxygen transport protein in A. ornata despite having a low degree of sequence homology with other hemoglobins (5–8) yet exhibits significant peroxidase activity approaching that of HRP even though it possesses neither structural nor sequence homology with any known peroxidase.

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1 Abbreviations: DHP, dehaloperoxidase; Hb, hemoglobin; HRP, horseradish peroxidase; Mb, myoglobin; 4IP, 4-iodophenol; 4BP, 4-bromophenol; DCP, 2,4-dichlorophenol; TBP, 2,4,6-trichlorophenol; TCP, 2,4,6-trichlorophenol; TFP, 2,4,6-trifluorophenol; X, trihalophenol; DXQ, dihalophenol; RFQ-CW-EPR, rapid-freeze-quench continuous wave electron paramagnetic resonance spectroscopy; Compound I, two-electron-oxidized heme species when compared to the ferric form, commonly as an $\text{Fe}^{\text{IV}}=\text{O}$ porphyrin $\pi$-cation radical; Compound II, one-electron-oxidized heme species when compared to the ferric form, commonly as an $\text{Fe}^{\text{III}}=\text{O}$ or $\text{Fe}^{\text{IV}}=\text{OH}$; Compound III, oxyferrous ($\text{Fe}^{\text{IV}}=\text{O}$; $\text{Fe}^{\text{III}}=\text{O}_2$) state of the enzyme; Compound ES, two-electron-oxidized state containing both a ferryl center ($\text{Fe}^{\text{IV}}=\text{O}$) and an amino acid (tryptophan or tyrosyl) radical, analogous to Compound ES in cytochrome c peroxidase; Compound RH, “reversible heme” state of dehaloperoxidase, formed from the decay of Compound ES in the absence of substrate.
DHP with its physiological oxidant, H$_2$O$_2$, yields what has been characterized to date as Compound II (7), which is formally a one-electron-oxidized heme center. Thus, the question of how DHP oxidizes its phenol substrates by two electrons when it appears to form only a one-electron oxidized intermediate needs to be addressed.

Several possible mechanisms for a net two-electron oxidation exist, including the simplest notion that two sequential one-electron oxidations are required for DHP to oxidize a trihalophenol, the first generating a trihalophenoxy radical intermediate and the second yielding a trihalocyclo-hexadienone cation which reacts with a solvent water molecule to yield the dihaloquinone product. These intermediates have been proposed by Osborne et al. to form during the oxidation of trichlorophenol to dichloroquinone by horse heart myoglobin (3). Another option is that DHP oxidizes trihalophenols by only one electron, forming a trihalophenoxy radical intermediate, which can undergo disproportionation with a second radical species, giving both trihalophenol and dihaloquinone. This proposed mechanism is reminiscent of ascorbate peroxidase (APX), which has been shown to oxidize ascorbate via a one-electron process to monohydroascorbate, the latter undergoing disproportionation to yield ascorbate and dehydroascorbate (13). A third possibility is that the intermediate in DHP formed from the reaction of the ferric enzyme with hydrogen peroxide is electronically similar to that of Compound ES in CcP, which has been shown to be isoelectronic with Compound I but in addition to the ferryl moiety also possesses a protein radical as the second oxidizing equivalent rather than a porphyrin-based one. This may be the favored mechanism, as it would allow for product formation via either (i) a direct two-electron oxidation (direct formation of the trihalocyclohexadienone cation) without generating radical intermediates (which could be potentially harmful if formed in the coelom of A. ornata) or (ii) two sequential one-electron oxidations via a transiently formed trihalophenoxy radical that would be further oxidized to the diene cation (3), potentially without first diffusing out of the active site pocket.

To explore mechanistic hypotheses, we have undertaken a comprehensive UV-visible and electron paramagnetic spectroscopic study of the DHP intermediate formed from the reaction of the ferric enzyme with hydrogen peroxide under a variety of conditions using both stopped-flow and rapid-freeze-quench methods and have determined this intermediate to be DHP Compound ES. We have further explored the stability of this Compound ES and demonstrated the existence of a competitive pathway between product formation and decay to a novel species, termed Compound RH, which possesses attenuated levels of dehaloperoxidase activity. Our experimental design reveals mechanistic insights and kinetic descriptions of the intermediates in DHP, which have not been previously reported. We propose an updated catalytic cycle which provides a clearer understanding of the link between peroxidase and oxygen transport activities unique to this bifunctional enzyme.

**MATERIALS AND METHODS**

Buffer salts and acetonitrile (HPLC-grade) were purchased from Fisher Scientific. All other reagents and biochemicals, unless otherwise specified, were of the highest grade available from Sigma-Aldrich. Solutions of 2,4,6-trichlorophenol (TCP) were freshly prepared daily in 100 mM potassium phosphate (KP) buffer (variable pH) and kept at 4 °C. UV-visible spectra were recorded periodically to ensure that the TCP substrate had not degraded (molar absorptivity listed in Table S1 of the Supporting Information). Hydrogen peroxide solutions were also freshly made prior to each experiment. Initially, a 10 mM stock solution of H$_2$O$_2$ was prepared and maintained at 4 °C (typically less than 15 min), during which all other protein/substrate solutions were loaded into the stopped-flow apparatus. When prepared in this manner, the stock H$_2$O$_2$ solution did not exhibit any degradation over this time period as determined by UV-visible spectroscopic analysis of the hydrogen peroxide absorbance at 240 nm ($\varepsilon_{240} = 43.6$ M$^{-1}$ cm$^{-1}$) (18). The stock H$_2$O$_2$ solution was then diluted to the appropriate premixing concentration and immediately loaded into the stopped-flow apparatus. Wild-type (WT) DHP (six-His-tagged protein) expression and purification procedures were preformed as previously described (4). TCP (4.05 mM maximum solubility) (14) was employed throughout this study due to the low solubility (0.355 mM) of TBP in aqueous solution (15).

**Spectroscopic Studies.** Optical spectra were recorded on either an Agilent 8453 diode array spectrophotometer or a Cary 50 UV-visible spectrophotometer, both equipped with thermostated cell holders at 25 °C. The protoheme content was measured by the pyridine hemochrome assay using a $\Delta\varepsilon_{555}$ of 20.7 M$^{-1}$ cm$^{-1}$ (reduced − oxidized) for iron protoporphyrin IX (16, 17).

**Enzyme Assays.** All measurements were performed in octiplet using a SpectraMax Plus384 UV-visible plate reader equipped with 96-well plates. Assays were carried out at 25 °C in 100 mM KP buffer (pH 7.5) containing 5 mM EDTA (total volume of 200 µL). Catalase activity was measured spectrophotometrically by following the decrease over 60 s (linear least-squares fittings) of the hydrogen peroxide concentration (1.5, 30, and 60 mM) at 240 nm ($\varepsilon_{240} = 43.6$ M$^{-1}$ cm$^{-1}$), in the presence of 1 µM DHP (18).

**Preparation of Ferric DHP.** DHP was incubated with a 1.7-fold molar excess of potassium ferricyanide for 2 min at room temperature. Excess ferricyanide was removed using a Sephadex G-25 gel-filtration column. The protein was concentrated using an Amicon Ultra centrifugal filter equipped with a 10000 kDa molecular mass membrane, and the purity of DHP was determined as previously described (4). Only DHP samples that exhibited Reinheitzahl values (RZ) greater than 4.0 were utilized in this study. The concentration of DHP was determined spectrophotometrically ($\varepsilon_{406} = 116.4$ M$^{-1}$ cm$^{-1}$) (19).

**Stopped-Flow UV−Visible Spectrophotometric Studies.** Experiments were performed on a Bio-Logic SFM-400 triple-mixing stopped-flow instrument equipped with a diode array UV-visible spectrophotometer and were carried out at 20 °C in 100 mM KP buffer (pH 5.0 and 7.0). A constant temperature was maintained using a circulating water bath. Data were collected (900 scans total) over a three-time domain regime (2.5, 25, and 250 ms; 300 scans each) using Bio Kinet32 (Bio-Logic). Experiments were performed in double-mixing mode using an aging line prior to the second mixing step. The design of the experiments allowed for the mixing of DHP with either TCP or H$_2$O$_2$ followed by an aging time of 1.5, 30, or 60 s, followed by the second mix...
with the remaining (co-) substrate: (i) DHP + TCP → delay → + H2O2 or (ii) DHP + H2O2 → delay → + TCP. Concentrations after mixing were as follows: [DHP]f = 10 µM, [H2O2]f = 100 µM, and [TCP]f = 300 µM. All data were evaluated using Specfit Global Analysis System (Spectrum Software Associates) as pseudo-first-order reactions and fit with SVD analysis from one to three exponential curves where applicable. Kinetic data were baseline corrected using the Specfit autozero function.

Preparation of Reaction Intermediates by Freeze-Quench Methods. Rapid-freeze-quench experiments were performed with a BioLogic SFM 400 freeze-quench apparatus by mixing a 50 µM enzyme solution (final concentration) with a 10-fold excess of H2O2 solution in 100 mM potassium phosphate buffer (pH 5 and 7) at 25 °C. Reaction times were as follows: pH 5, 100 ms, 400 ms, 3.6 s, 36 s, and 60 s; and pH 7, 100 ms, 500 ms, 800 ms, 2 s, 12 s, and 60 s. A standard 4 mm outside diameter quartz EPR tube was connected to a Teflon funnel, and both the tube and the funnel were completely immersed in an isopentane bath at −110 °C. The reactions were quenched by spraying the mixtures into the cold isopentane, and the frozen material so obtained was packed at the bottom of the quartz tube using a packing rod equipped with a Teflon plunger. Samples were then transferred to a liquid nitrogen storage dewar until they were analyzed.

X-Band EPR Spectroscopy. EPR spectra were recorded with an X-band (9 GHz) Varian E-9 EPR spectrometer (Varian, El Palo, CA). A standard 3 mm × 4 mm quartz EPR tube was filled with a sample and placed into a quartz finger dewar insert filled with liquid nitrogen. The temperature of the samples was kept at 77 K for the duration of the data acquisition, which required periodic refilling of the dewar due to the evaporation of the liquid nitrogen during longer acquisition runs. The typical spectrometer settings were as follows: field sweep, 200 G; scan rate, 3.33 G/s; modulation frequency, 100 kHz; modulation amplitude, 4.0 G; and microwave power, 2 mW. The exact resonant frequency of each EPR experiment was measured by an EIP-578 (PhaseMatrix, San Jose, CA) in-line microwave frequency counter and is indicated in the figure captions. Typically, 20 and 200 individual scans were averaged to achieve sufficient signal to noise for the spectra obtained at short quench and long quench times, respectively.

RESULTS

UV–Visible Spectroscopic Studies of Ferric DHP in the Presence of TCP. The electronic absorption spectra of ferric metahaem DHP at pH 5 and 7 are presented in Figure 1. TCP exhibits absorbance maxima at 285 and 311 nm at pH 5.0 (Figure 1A) and pH 7.0 (Figure 1B), respectively. Although there is evidence for binding of substrate at an internal site (8), there is also evidence for an external (20, 21) binding site. The spectra in Figure 1 show that TCP has a minimal effect on the heme spectra, and it is believed that TCP binds at an external binding site in the stopped-flow studies reported here.

Stopped-Flow UV–Visible Characterization of Compound ES in DHP. Single-mixing stopped-flow UV–visible spectroscopic methods were employed to detect DHP Compound ES. At pH 7, when a solution of ferric DHP [UV–visible spectrum, 407 (Soret), 504, 538, 635 nm] was rapidly mixed (2 ms) with H2O2, a transient species was observed (Figure 2 and Table 1), the spectral features of which [UV–visible, 420 (Soret), 554, 585 nm] we ascribe to a ferryl-containing DHP intermediate based upon previous characterization (4, 7, 22) and comparison to other known Fe(IV)–oxo species-containing hemoproteins (11, 12). As the ferryl intermediate of Compound ES is likely indistinguishable from that of Compound II by UV–visible spectroscopy, we assign this intermediate here as DHP Compound ES on the basis of these results and those of our EPR spectroscopic study (vide infra). Values of kobs for formation of this new species were linearly dependent on H2O2 concentration (2.5–10-fold excess per heme), giving a bimolecular rate constant of (3.56 ± 0.02) × 104 M−1 s−1. Under these conditions, and in the absence of substrate, DHP Compound ES decays to a stable species [UV–visible, 411 (Soret), 530, 564 nm; kobs = 0.0167 ± 0.0003 s−1], which we have termed Compound RH.

Similar reactivity is observed at pH 5. Ferric DHP [UV–visible, 405 (Soret), 504, 538, 636 nm] is converted to Compound ES [UV–visible, 419 (Soret), 545, 585 nm; kobs = (2.78 ± 0.01) × 104 M−1 s−1], which further decays to Compound RH [UV–visible, 410 (Soret), 530, 590 nm; kobs = 0.0701 ± 0.0001 s−1] (Figure S1 of the Supporting Information). Hence, the rate of Compound RH formation is ~4 times greater at pH 5 than at pH 7.

Reaction of Preformed Compound ES with TCP Substrate. Stopped-flow UV–visible spectroscopy was employed to monitor the reaction between preformed DHP Compound ES and TCP. In a double-mixing experiment, 10 µM DHP was first reacted with a 10-fold excess of H2O2, allowed to incubate for 1.5 or 0.9 s, corresponding to the maximum accumulation of DHP Compound ES at pH 7 or 5, respectively, and then subsequently mixed with a 30-fold excess of TCP, which resulted in the regeneration of ferric (resting) DHP (Figure 3 and Figure S2 of the Supporting Information).
Under these conditions, nearly identical quantities of the DCQ product are formed at both pH values (Figure 4). However, it is interesting to note that at pH 5.0, the disappearance of Compound ES and the return of the enzyme to its resting state (8.47 ± 0.05 s⁻¹) are not directly coupled with product formation, which occurs subsequent to that process (0.13 ± 0.01 s⁻¹). On the other hand, at pH 7.0 the disappearance of DHP Compound ES (0.11 s⁻¹) is concomitant with the appearance of product (0.14 s⁻¹), although this observation does not prove that they are coupled. Nonenzymatic control experiments confirmed the necessity for DHP at both pH values, as no DCQ product was observed in the absence of the enzyme.

When the double-mixing experiments described above were repeated with longer incubation times that permitted Compound ES to completely decay to Compound RH prior to mixing with TCP, neither substrate loss (311 nm) nor product formation (275 nm) was observed during the 85 s time scale of these stopped-flow experiments. Specifically, at pH 5, Compound RH is nearly fully formed after 30 s (Figure S1c of the Supporting Information), with little to no Compound ES present, and neither the 30 nor 60 s incubation time exhibited product formation at this pH (Figure 4), indicating that Compound RH possesses a much lower catalytic activity than ferric DHP (vide infra). At pH 7, a 30 s incubation time results in a partial mixture of Compounds ES and RH being observed (Figure 2C), concomitant with partial substrate oxidation, whereas the 60 s incubation period led to no product formation, consistent with the nearly complete decay of Compound ES to Compound RH observed in the single-mixing experiments during this time frame. Thus, the extent of product formation is directly correlated with the amount of Compound ES present, strongly indicative that this intermediate is an active oxidant in DHP.

In Situ Compound ES Formation in the Presence of TCP. In contrast to the experiments described above in which preformed Compound ES was reacted with TCP, we employed double-mixing stopped-flow UV-visible spectroscopic methods to examine if preincubation of ferric DHP with TCP, followed by the addition of a 10-fold excess of H₂O₂, led to the formation of DCQ product (275 nm) via a transiently formed Compound ES intermediate. Under these conditions, the yield of DCQ product was nearly identical with that found for the reaction of fully preformed Compound ES with TCP (vide supra), at both pH 7 (Figure 5) and pH 5 (Figure S3 of the Supporting Information). When TCP was preincubated with DHP, there was no dependence on the delay time (1.5—60 s) prior to addition of H₂O₂ (Figure 6 and Figure S4 of the Supporting Information for pH 7 and 5, respectively). DCQ product formation had pseudo-first-order constants (kobs) of 0.11 and 0.22 s⁻¹ for pH 7.0 and 5.0, respectively. DCQ was also observed to be unstable under these conditions, undergoing side reaction(s) leading to its slow loss at longer time periods which were more apparent at pH 5 than at pH 7, but this chemistry was not further explored.

We also did note a pH-dependent difference with respect to the heme species observed. At pH 7.0, Compound ES [UV-visible, 420 (Soret), 545, 585 nm] is distinctly formed, with approximately the same rate (kobs = 3.11 ± 0.02) × 10⁴ M⁻¹ s⁻¹ as when TCP was absent [kobs = 3.56 ± 0.02] × 10⁴ M⁻¹ s⁻¹], and returns to the ferric state (UV-visible, 407, 504, 538, 578, 633 nm) with approximately the same rate (kobs = 0.12 s⁻¹) as product formation (0.11 s⁻¹). At pH 5.0, no distinct Compound ES spectrum is observed upon reaction of hydrogen peroxide with DHP incubated in the presence of TCP; the UV-visible features [406 (Soret), 504, 538, and 635 nm] match those of the resting (ferric) enzyme, with the caveat that a minor broadening on the red side may be indicative of Compound ES formation and disappearance within the stopped-flow mixing time (2 ms).
Characterization and Reactivity of DHP Compound ES

Rapid-freeze-quench methods were employed to stabilize intermediates of the reaction between DHP (final concentration of 50 μM) and a 10-fold excess of hydrogen peroxide at both pH 7 and 5 for consequent characterization by continuous wave (CW) EPR. X-Band CW EPR spectra of DHP samples recorded at pH 7.0 with various quench times are shown in Figure 7. The shapes of all the EPR spectra measured from the samples collected with quenching times of 500 ms, 800 ms, and 2 s were found to be identical, the only difference being in the signal intensities. The maximal concentration of the radical is observed over the period of Compound ES formation (Figure 2C). The position of the signal is characterized by an average g factor of 2.0058. The shape of the signal is best described by an anisotropic quintet. On the basis of the signal g factor and a partially resolved hyperfine structure with a peak-to-peak line width of ~21 G, this EPR signal was assigned to a tyrosyl radical (23). Samples with longer incubation times of 12 and 60 s have similar average g factors (g ≈ 2.005) and the same peak-to-peak line width of 21 G; however, they do not exhibit resolved hyperfine structure and have much lower intensity [concomitant with the loss of the ferryl UV−visible spectrum in our component analysis (Figure 2C)].

The EPR spectrum of Compound ES was similarly recorded at various quench times at pH 5 (Figure 8). The spectrum is centered at g = 2.0058 and shows a partially resolved hyperfine splitting described as an “anisotropic septet” (29). A very weak shoulder is observed at g = 2.035 (Figure S5 of the Supporting Information) that could be an indication of the formation of a peroxyl radical since the samples were prepared under aerobic conditions (30), but the signal intensity is too low to warrant further speculation about its origin. At longer quench times when the component analysis indicates little to no remaining Compound ES by UV−visible spectroscopy (Figure S1c of the Supporting Information), the line shape of the EPR signal is drastically different, its signal intensity has dropped considerably, and the hyperfine splitting features are lost. Specifically, the signal has a very broad spectral feature in the g ≈ 2.04 region and sharper features at g = 2.0085 and g = 1.995.

Unfortunately, the low g factor spectral resolution of these CW X-band experiments does not permit unambiguous identification of the radical species on the basis of magnetic parameters alone. This ambiguity could be resolved by high-field (95 GHz) EPR experiments coupled with mutagenesis studies that are planned for the near future.

Formation and Reactivity of Compound RH. As described above, in the absence of reducing substrate, a new, unique species of DHP, termed Compound RH to denote that it is a reversible heme intermediate, is formed upon the decay of Compound ES. Compound RH was found to be robust to other sample preparation procedures. While its UV−visible features remained constant, we did observe that some protein

<table>
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<th>Compound</th>
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<th>pH 5</th>
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<tr>
<td></td>
<td>λ_{max} (nm)</td>
<td>k_{obs} (M^{−1} s^{−1})</td>
</tr>
<tr>
<td>ferric</td>
<td>407, 504, 538, 635</td>
<td>not applicable</td>
</tr>
<tr>
<td>Compound ES</td>
<td>420, 545, 585</td>
<td>(3.56 ± 0.02) × 10^{4} M^{−1} s^{−1}</td>
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<tr>
<td>Compound RH</td>
<td>411, 530, 564</td>
<td>0.0167 ± 0.0003 s^{−1}</td>
</tr>
<tr>
<td>Compound III</td>
<td>417, 542, 578</td>
<td>not determined</td>
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Table 1: UV−Visible Spectroscopic Data and Kinetic Parameters for the Oxidized Intermediates of DHP
precipitation occurred when the long-term (several hours) room temperature stability of the Compound RH species was investigated. Addition of excess potassium cyanide (600 mM) to Compound RH resulted in a new UV-visible spectrum observed \(421\) (Soret), \(545, 575\) (sh) (data not shown), consistent with the formation of a DHP ferricyanide adduct \((31)\), providing further evidence that Compound RH is a single species.

To test the reactivity of the Compound RH species, single-mixing stopped-flow UV-visible spectroscopy was used to monitor the reaction between Compound RH (10 \(\mu\)M) and either a 10- or 100-fold molar excess of \(\text{H}_2\text{O}_2\). In both cases, no reaction was observed [both pH 5 and 7 were investigated (data not shown)]. In spite of its inability to form high-valent iron–oxo intermediates, Compound RH still possesses attenuated levels of dehaloperoxidase activity, exhibiting a 6-fold lower reactivity with TCP as a substrate when compared to ferric DHP.

While Compound RH displays a lack of reactivity with \(\text{H}_2\text{O}_2\) by stopped-flow UV-visible spectroscopy, it is readily reduced with sodium dithionite at either pH 5 or 7 to yield ferrous (deoxy) DHP [UV-visible, \(432\) (Soret), \(557, 626\) nm], which upon exposure to dioxygen led to formation of oxyferrous DHP [UV-visible, \(417\) (Soret), \(542, 578\) nm]. Further oxidation with potassium ferricyanide, followed by
Characterization and Reactivity of DHP Compound ES

DHP has been shown to perform the net two-electron oxidation of phenol (X−A−OH + H2O2 → O═A═O + HX + H2O) to yield the corresponding quinone, consistent with the reactive species being a two-electron-oxidized intermediate such as Compound I or Compound ES (10−12). However, the reaction of DHP with its physiological oxidant, H2O2, yields what had been characterized to date as Compound II, which is formally a one-electron-oxidized heme center. Thus, the question of how DHP oxidizes its phenol substrates by two electrons when it has been shown to only form a one-electron-oxidized intermediate needs to be addressed. This study provides further characterization of the intermediate formed from the reaction of ferric dehaloperoxidase with hydrogen peroxide, which suggests that this species is similar to the two-electron-oxidized Compound ES of CcP.

We employed stopped-flow UV−visible spectroscopic methods to monitor the reaction of ferric DHP with its physiological oxidant, H2O2. The first intermediate observed had spectral features which unequivocally match those for a ferryl-containing species lacking a porphyrin π-cation radical, such as Compound ES or Compound II. This implies either that Compound I is not formed by this reaction or more likely that it is formed transiently but undergoes rapid reduction to the observed Compound ES intermediate (vide infra). Interestingly, in the absence of a reducing substrate, the Compound ES intermediate was found to be unstable and converted to a new, stable species which we have termed Compound RH on the basis of the fact that it is a reversibly formed heme intermediate (RH, reversible heme). In the presence of substrate, however, Compound ES returned to the resting state, concomitant with formation of the quinone product. We examined this reaction with TCP using both preformed and in situ-generated Compound ES, and in both cases, the amount and rate of DCQ product generated were identical and independent of pH. The implication is that Compound ES is likely the actual species which is catalyzing TCP oxidation, rather than Compound I. If this were not the case, then one might expect different results between the amount and rate of product formation for in situ versus preformed Compound ES reactions. While this does not absolutely preclude a transiently formed Compound I species from oxidizing TCP, it strongly suggests that Compound ES is the active oxidant. Dichloroquinone product formation may result from either a single two-electron oxidation of the trichlorophenol substrate by Compound ES or two sequential one-electron oxidations, both pathways being indistinguishable under the multiple-turnover conditions examined in this study.

Stopped-flow UV−visible and rapid-freeze-quench EPR spectroscopic studies were employed to follow the formation of the high-valent iron(IV)−oxo and protein radical species, respectively, for the reaction of DHP with H2O2 leading to Compound ES. The UV−visible spectroscopic data indicate the direct formation of a classical ferryl-containing species, similar to that formed in HRP and Mb. Moreover, a recent ENDOR study by Hoffman and co-workers (32) supports the assignment as FeIV=O, and not the protonated analogue FeIV−OH (as in chloroperoxidase). The lack of an observable Compound I species suggested the presence of an endogenous reducing agent. As the crystal structure of DHP does not show any other redox capable cofactor (e.g., flavin, another metal center, etc.), we investigated by RFQ-CW-EPR spectroscopy whether a protein side chain could be responsible for the rapid reduction of a transiently formed Compound I species to the observed Compound ES intermediate, resulting in the formation of a protein radical. Indeed, we have identified the presence of a protein radical whose maximal concentration is observed over the period of Compound ES formation identified from UV−visible spectroscopy. As a result, we present here for the first time clear evidence of DHP Compound ES as a two-electron-oxidized intermediate, with one oxidation equivalent centered on the ferryl moiety and the second oxidation equivalent located on a protein side chain as a radical. Thus, DHP Compound ES bears a strong resemblance to CcP Compound ES, which has been extensively studied and shown to possess both a ferryl heme center and a radical on Trp101.

Analysis of the signal g factor and the partially resolved hyperfine structure present in our EPR data, with a peak-to-peak line width of ∼21 G, suggests that the radical in DHP Compound ES likely resides on a tyrosine residue initially (23). The longer incubation times of 12 and 60 s exhibit a similar average g factor (g ≈ 2.005) and the same peak-to-peak line width of 21 G; however, they do not show resolved hyperfine structure and have much lower intensity [concomitant with the loss of the ferryl UV−visible spectrum in our component analysis (Figure 2C)]. Thus, it is difficult to comment extensively on the origin of these signals observed at later times. These spectra could be attributed to a different Tyr-based radical than the one initially observed, or the signal could also be an admixture of Tyr- and Trp-based radicals (24). Another option to consider is a radical originating from a cysteine residue, which is usually characterized by high g factor anisotropy that would be resolved in the X-band spectra (25−27). Although Cys-based radicals have been accepted as intermediates in metalloprotein cycles, they are not commonly observed by freeze-quench EPR.

![Figure 8: EPR spectra of the radical(s) in DHP Compound ES at pH 5.](image-url)
spectroscopy. However, Cys-based radicals should not be completely ruled out as intermediates at the quenching times of 12 and 60 s as they have been shown to form in human Mb under similar conditions (28).

Of the five tyrosine residues which DHP possesses [Tyr16, Tyr28, Tyr34, Tyr38, and Tyr107 (Figure 9)], only two are reasonably close to the heme prosthetic group to be considered likely candidates for reducing Compound I initially, Tyr34 and Tyr38; the closest contact between Tyr34 and the heme edge is 5.56 Å (Tyr C with Cys73 (16.29 Å) contacts are similarly distant, making them unlikely candidates for the residues responsible for the initial reduction of the transiently formed Compound I intermediate. Thus, on the basis of both the available structural studies of DHP and our spectroscopic analysis presented here, we suggest that the initial site of radical formation in DHP Compound ES is either Tyr34 or Tyr38.

While the definitive identification of the specific residue that gives rise to the initial protein radical in DHP Compound ES will be the subject of future high-field EPR investigations coupled with mutagenesis studies, the time-dependent changes observed in the protein radical signal suggest either a change in the local electronic structure of the radical or a migration of the radical out of the active site to other redox active protein side chains upon decay of Compound ES. This latter conjecture is interesting, given that the reaction of sperm whale Mb with hydrogen peroxide yields covalent dimers that arise from the coupling of surface tyrosyl radicals (Tyr151), which results from a radical migration out of the Mb active site (33). Similarly, it has been demonstrated that human Mb also forms a covalent dimer (34), but through surface cysteines (Cys110) forming a disulfide link, consistent with our putative radical observed at long quench times at pH 5 possessing a similar radical migration pathway, with surface radicals leading to the oxidation of TCP that binds to DHP through a hypothesized external binding pocket. This would not be unlike that of CcP, whose initial Trp191 radical in Compound ES leads to a radical migration which is ultimately responsible for oxidizing substrate (cytochrome c) at the surface of the peroxidase (protein–protein interface) (35). Thus, as CcP may have evolved from traditional (heme edge electron transfer) peroxidases an external binding interface for oxidizing cytochrome c, DHP may also similarly have evolved from Mb an external binding pocket for oxidizing trihalophenols. In accord with this hypothesis, tyrosine has been proposed to play a role in the peroxidase mechanism of Mb (36). This hypothesis will be particularly interesting if our supposition is that Tyr34 is the site of the radical in DHP Compound ES, as this residue is located at the surface of DHP yet is also in the proximity of the heme (Figure 9), and thus could serve as a redox conduit between the hypothesized external binding pocket and the active site.

From the initial observation in the X-ray crystal structure that 4-iodophenol (4IP) binds at an internal site in the distal pocket of DHP (8), the possibility of substrate binding to both internal and external binding pockets has been investigated with a number of techniques, including Fourier transform infrared (FTIR) (20), electron paramagnetic resonance (EPR) (37), nuclear magnetic resonance (NMR) (21), and resonance Raman experiments (Thompson and S. Franzen, unpublished results). Solution studies (21) and cryogenic studies (20, 37) have been carried out using 2,4,6-trifluorophenol (TFP) as a model of the native substrate 2,4,6-tribromophenol (TBP), which is relatively insoluble. EPR and HYSCORE experiments (37) are consistent with a change from six-coordinate high-spin to five-coordinate high-spin upon TFP binding at pH 6.0. TFP was used in those experiments because of its relatively high solubility, which facilitates detection under the conditions of the EPR experiment (4 K). FTIR experiments conducted on the carbonmonoxy form of DHP (DHP–CO) at cryogenic temperature (20) show that there is a large effect of TFP binding on the CO stretching frequency and rebinding kinetics at pH 5.5, but not at pH 7.0. However, near ambient temperature and physiological pH, TFP does not bind to DHP–CO (20). On the basis of the binding of 4IP in the distal pocket of DHP in the X-ray crystal structure, one hypothesis is that TFP is bound in the distal pocket of DHP–CO at low pH and at low temperatures. Hyperfine 1H NMR experiments show that 4-bromophenol (4BP) and 2,4-dichlorophenol (DCP) interact with the metcyano form (DHP–CN) at ambient temperature through changes in the position of the internal heme edge Phe201 residue and the 3-CH3 heme methyl (21). TFP does not produce these changes at ambient temperature, but there are shifts in 19F resonances of the substrate when it interacts with DHP, which can be detected by 19F NMR. From these data, one can further hypothesize that there is an external binding site (21) as well as the internal binding site observed by X-ray crystallography (8). Further confirmation can be found in resonance Raman experiments on 4IP and 4BP, which also show a change in core size marker modes when substrate binds (S. Franzen and Thompson, unpublished results). These experiments confirm that there is a change from six-coordinate high-spin in the resting metaperoxo form of DHP to five-coordinate high-spin when 4IP and 4BP bind. Thus, while 4IP and 4BP appear to bind inside the distal

**FIGURE 9: Active site of DHP showing all tyrosine residues present in the enzyme.** The proximal and distal histidines, His89 and His55, respectively, are provided for orientation. Coordinates (2QFK) were obtained from the Protein Data Bank and displayed using Pymol.
Scheme 2: Proposed Catalytic Cycle for Dehaloperoxidase

In the absence of substrate, we observed that DHP Compound ES forms a new, stable species, termed Compound RH, which is unreactive toward \( \text{H}_2\text{O}_2 \) and, as a consequence, possesses attenuated levels of dehaloperoxidase activity. While peroxidase inactivation is a known phenomenon in HRPs or CcPs, where heme bleaching is the primary culprit (38), Compound RH is still active and can reversibly return to the deoxy ferrous state under mild reduction conditions. Furthermore, Compound RH has not been reported in myoglobin or hemoglobin previously, and is therefore unique to DHP. Compound RH is still active and can reversibly return to the deoxy ferrous state under mild reduction conditions. Furthermore, Compound RH has not been reported in myoglobin or hemoglobin previously, and is therefore unique to DHP (33, 39–42). Thus, as it is not found in either the monofunctional peroxidases or the \( \text{O}_2 \) transport globins, DHP Compound RH represents a novel species which may be a result of the bifunctional nature of DHP: the inactivated (dehalo)peroxidase can be regenerated back to an active oxygen transport protein via a reduction pathway that is specific for globins, but not peroxidases. When DHP is functioning as a peroxidase, its inactivation is necessary to prevent nonspecific oxidation of other metabolites from occurring when trihalophenol substrate is absent. However, while heme bleaching is a normal route for peroxidase inactivation, this could be metabolically costly for DHP, given that it is the hemoglobin in \( A. \ ornata \). Thus, formation of Compound RH allows dehaloperoxidase to be inactivated without sacrificing the protein to heme bleaching while at the same time allowing for its functional switch back to an oxygen transport protein upon reduction. While the exact nature of the Compound RH species will require further study, it nevertheless represents an interesting and novel observation in this study.

A catalytic cycle can be proposed on the basis of the reversible formation of Compound RH. The catalytic cycle shown in Scheme 2 confers “hydrogen peroxide reductase” activity to dehaloperoxidase. The use of reducing agents to scavenge reactive oxygen species enzymatically is not unprecedented, with members of the peroxiredoxin family (Prx, also termed the thioredoxin peroxidases and alkyl-hydroperoxide-reductase-C22 proteins) and superoxide reductase (SOR) being perhaps the most well-known examples for \( \text{H}_2\text{O}_2 \) and superoxide reduction, respectively (43, 44). While disproportionation of \( \text{H}_2\text{O}_2 \) may be preferable over its reduction from a metabolic viewpoint, DHP does not exhibit significant catalase activity, although myoglobin has been shown to consume (noncatalytic) up to 8 equiv of \( \text{H}_2\text{O}_2 \) (33, 41, 42). Overall, should a partner reductase similar to Mb reductase be identified for DHP, the door for DHP to be considered a trifunctional protein capable of peroxidase, \( \text{O}_2 \) transport, and \( \text{H}_2\text{O}_2 \) scavenging activities would be opened.

On the basis of the results obtained from these stopped-flow UV—visible and RFQ-EPR spectroscopic experiments, and through modification of previously established mechanisms for the general function of peroxidases (45), we propose the following catalytic cycle for the in vitro peroxide-dependent oxidation of ferric DHP from \( A. \ ornata \) in the presence and absence of trihalophenol (Scheme 2). Ferric DHP reacts with 1 equiv of \( \text{H}_2\text{O}_2 \), transiently forming Compound I (step i), which then undergoes rapid endogenous electron transfer to generate the observed Compound ES intermediate and protein radical (step ii). A bifurcation in the mechanism occurs which is dependent upon substrate. In the presence of trihalophenol, DHP Compound ES is reduced by two electrons, thereby regenerating the ferric state of the enzyme, and forming the dihaloquinone product (step iii-a). In the absence of substrate, however, Compound RH is formed (iii-b) by a not-yet-understood process and can subsequently be reduced to the ferrous enzyme (iv) and bind dioxygen to form the oxyferrous intermediate (v). Autoxidation of oxyferrous DHP leads to the formation of the ferric enzyme (vi). The existence, and by extension identity, of a possible sixth ligand in Compound RH is unknown at this time, and this ambiguity is represented by the bound X in Scheme 2. We tentatively assign the oxidation state of Compound RH as a ferric heme on the basis of the evidence that (i) Compound RH exhibits a “ferric-like” heme spectrum that matches neither the ferryl nor the ferrous spectra of DHP (Table 1), (ii) Compound RH forms from the decay of an iron(IV)–oxo species, which for most heme proteins yields a ferric enzyme, and (iii) Compound RH can be reduced to the ferrous enzyme, implying that it does not start as a ferric heme. Further studies will be necessary to definitively assign the oxidation state of the heme in Compound RH.

**CONCLUSION**

This study addresses a number of key questions pertaining to the nature and catalytic competency of the Compound ES intermediate in DHP. Our spectroscopic and biochemical characterization of DHP Compound ES suggests that this species is similar to the two-electron-oxidized Compound ES of CcP in that it possesses both a ferryl heme center and a protein radical. Furthermore, our results are consistent with Compound ES being an active species responsible for trihalophenol oxidation, as opposed to Compound I. The data support the hypothesis that there is an external substrate binding pocket in DHP, in which case the data reported here indicate that there may be a radical migration pathway in DHP analogous to that in CcP. A peroxidase-attenuated species unique to DHP, namely Compound RH, was also
identified, and a role for its formation as a protective species against unwanted oxidation chemistry was hypothesized. As Compound RH is unreactive toward further oxidation, it was found that reduction of Compound RH regenerates oxygenous DHP, and this suggests that the recovery of the oxygen transport function from an attenuated peroxidase species via reduction (possibly globin reductase) is a consequence of the bifunctional nature of this protein and may represent the chemical process that links the oxygen transport and peroxidase activities in dehaloperoxidase.

**SUPPORTING INFORMATION AVAILABLE**

Molar absorbances for 2,4,6-trichlorophenol at pH 5, 6, 7, and 7.5 (Table S1), stopped-flow UV–visible spectroscopic monitoring of the reaction between DHP and a 10-fold molar excess of H2O2 at pH 5.0 (Figure S1), stopped-flow UV–visible spectroscopic monitoring of the double-mixing reaction between preformed DHP Compound ES and TCP at pH 5 (Figure S2), stopped-flow UV–visible spectroscopic monitoring of the double-mixing reaction of ferric DHP preincubated with TCP for 1.5 s prior to its reaction with H2O2 at pH 5 (Figure S3), stopped-flow UV–visible spectroscopic monitoring of the double-mixing reaction of ferric DHP preincubated with TCP for 60 s prior to its reaction with H2O2 at pH 5 (Figure S4), and rapid-freeze-quench data at pH 5 (Figure S5). This material is available free of charge via the Internet at http://pubs.acs.org.

**REFERENCES**


