Pre-steady-state Kinetics Reveal the Substrate Specificity and Mechanism of Halide Oxidation of Truncated Human Peroxidasin 1*

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Human peroxidasin 1 is a homotrimeric multidomain peroxidase that is secreted to the extracellular matrix. The heme enzyme was shown to release hypohalous acid that mediates the formation of specific covalent sulfilimine bonds to reinforce collagen IV in basement membranes. Maturation by proteolytic cleavage is known to activate the enzyme. Here, we present the first multimixing stopped-flow study on a fully functional truncated variant of human peroxidasin 1 comprising four immunoglobulin-like domains and the catalytically active peroxidase domain. The kinetic data unravel the so far unknown substrate specificity and mechanism of halide oxidation of human peroxidasin 1. The heme enzyme is shown to follow the halogenation cycle that is induced by the rapid H2O2-mediated oxidation of the ferric enzyme to the redox intermediate compound I. We demonstrate that chloride cannot act as a two-electron donor of compound I, whereas thiocyanate, iodide, and bromide efficiently restore the ferric resting state. We present all relevant apparent bimolecular rate constants, the spectral signatures of the redox intermediates, and the standard reduction potential of the Fe(III)/Fe(II) couple, and we demonstrate that the prosthetic heme group is post-translationally modified and cross-linked with the protein. These structural features provide the basis of human peroxidasin 1 to act as an effective generator of hypohalous acid, which mediates the formation of covalent cross-links in collagen IV.

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Kinetics of Interconversion of Redox Intermediates of Pxd01

In this work, to study the reactivity of the peroxidase domain and the effect of other domains on catalysis, several truncated variants of hsPxd01 were expressed recombinantly in HEK cells (5). One monomeric construct composed of the four Ig domains and the peroxidase domain (hsPxd01-con4) (Fig. 1A) was superior in terms of yield, heme insertion, spectroscopic features, and catalytic activity. This allowed for the first time a comprehensive study of the kinetics of interconversion of the relevant redox intermediates of the halogenation cycle. Here, we report the standard reduction potential of the Fe(III)/Fe(II) couple of the peroxidase domain and apparent bimolecular rate constants for cyanide binding as well as the formation and reduction of compound I mediated by hydrogen peroxide, bromide, iodide, and thiocyanate at pH 7.4. Based on the available structural, kinetic, and thermodynamic data of the homologous human peroxidases, we provide a mechanism for the halogenation cycle. Here, we report the standard reduction potential of the Fe(III)/Fe(II) couple (9, 10).

Covalent attachment of the prosthetic group could also be confirmed by precipitation of the recombinant protein by acetone at pH 4.5 resulting in colorless supernatants (data not shown). Furthermore, the determined spectral and redox properties clearly suggested the establishment of heme–protein bonds (see below). It is well known that these post-translational modifications considerably influence both the spectral signatures of a heme protein in its ferric state and the standard reduction potential of the Fe(III)/Fe(II) couple (9, 10).

In this context, it is important to note that the purification protocol (see under “Experimental Procedures”) could be significantly improved by including a 48-h incubation step prior to buffer exchange and affinity chromatography. This period seemed to be necessary for establishment of covalent bonds. The modification of the prosthetic group is reflected by a gradual transition of the UV-visible spectrum of freshly purified hsPxd01-con4. Spectral transition included a red-shift of the Soret band from 410 to 412 nm together with establishment of Q-bands at 510, 547, and 590 nm and a charge transfer (CT) band at 637 nm (Fig. 1C, gray spectrum). This spectrum is reminiscent of that of ferric LPO (Fig. 1C, gray spectrum). The peroxidase domain of hsPxd01 has a high amino acid sequence homology with LPO (34% identity and 53% similarity) (5). In the known crystal structure of LPO, continuous electron densities confirmed the establishment of heme-protein bonds. The presence of two ester bonds between the modified prosthetic heme group and conserved Asp and Glu residues (7). Sequence alignment and modeling demonstrated that these acidic amino acids are fully conserved in hsPxd01 (i.e. Asp626 and Glu780) (3, 5).

Heme modification was also underlined by spectroelectrochemical studies on hsPxd01-con4. Fig. 2 shows a representative redox titration with fully oxidized and fully reduced hsPxd01-con4 depicted in green and brown, respectively. Upon reduction, the Soret peak of the ferric protein shifted from 412 to 435 nm (Fig. 2) very similar to 5-coordinated ferrous lac-
toperoxidase and eosinophil peroxidase (11). From the corresponding Nernst plot (Fig. 2B), the standard reduction potential ($E^0$) of the Fe(III)/Fe(II) couple was calculated to be $-0.128 \pm 0.006$ V (25 °C and pH 7.4).

Cyanide Binding—Before investigating the kinetics of compound I formation and reduction, we probed the accessibility of the heme cavity and the homogeneity of the architecture of the active site by monitoring the kinetics of cyanide binding. It has been demonstrated that the post-translational modification of the prosthetic group in peroxidases from the heme peroxidase-cyclooxygenase superfamily is often not fully established resulting in some structural heterogeneity of the heme cavity that can easily be probed by studying the kinetics of cyanide binding (7, 10). In the case of hsPxd01-con4, cyanide binding resulted in the transition of the high spin ($S = 5/2$) Fe(III) state to the low spin ($S = 1/2$) Fe(III) state (Soret maximum at 431 nm, bands at 558 and 588 nm, and loss of the CT band at 637 nm) with clear isosbestic points at 360 and 443 nm, respectively.

The kinetics of hsPxd01-con4 oxidation mediated by H$_2$O$_2$ was followed by the decrease of absorbance at 412 nm. The reaction was biphasic with a dominating rapid first phase ($>80\%$ of $A_{412}$ nm). The corresponding pseudo first-order rate constants $k_{obs(1)}$ and $k_{obs(2)}$ were obtained from double exponential fits. From the slope of the plots of the respective $k_{obs}$ values versus hydrogen peroxide concentration, the apparent bimolecular rate constants for the dominating phase $k_{app(1)}$ ($1.8 \pm 0.1 \times 10^7$ M$^{-1}$ s$^{-1}$) and the second phase $k_{app(2)}$ ($6.9 \pm 0.6 \times 10^5$ M$^{-1}$ s$^{-1}$) (data not shown), were calculated, pH 7.4.

Furthermore, we could demonstrate that the rate of compound I formation was invariant within the pH range of 5.0–9.0 (Fig. 4D). A pK$_a$ value of 4.7 and a pK$_b$ value of 9.0 were calculated from the fit $k_{app} = a/(1 + x/b) \times (1 + c/x)$ with $a$ representing $k_{int}$, $x$ is the concentration of H$^+$; $b$ is pK$_a$, and $c$ is
The first inflection point might reflect the pK_a of the distal histidine that acts as proton acceptor in compound I formation of heme peroxidases (7), whereas the second one could be related with the alkaline transition of hsPxd01-con4. The UV-visible spectrum of the ferric high spin protein is invariant between pH 5 and 8.0 (5) but converts to a low spin spectrum with red-shifted Soret band (430 nm) at pH values 8.5 most probably reflecting the formation of a low spin hydroxide complex.

Addition of excess hydrogen peroxide to ferric hsPxd01-con4 converts the enzyme from the ferric state via compound I to compound II (oxoiron(IV) species) and, finally, to compound III, which resembles electronically oxymyoglobin or oxyhemoglobin (i.e. Fe(II)-O_2 ↔ Fe(III)-O_2^-) (Fig. 5). When hsPxd01-con4 was mixed with a 50-fold molar excess of hydrogen peroxide, the formation of predominantly compound II was observed with a heme Soret maximum at 432 nm and a broad Q band at 535 nm with a shoulder at 564 nm (Fig. 5, blue spectrum). With a 1000-fold molar excess of H_2O_2, compound II was converted to compound III resulting in a distinct UV-visible spectrum with a heme Soret maximum at 425 nm and prominent Q-bands at 552 and 588 nm (Fig. 5, cyan spectrum).

It was interesting to see that in contrast to LPO (14) but similar to myeloperoxidase (7), the conversion of compound I
to compound II was also dependent on the hydrogen peroxide concentration (Fig. 6). After incubating hsPxd01-con4 with equimolar H₂O₂ in the aging loop of the stopped-flow instrument for 200 ms, formed compound I was mixed with increasing concentrations of H₂O₂. Formation of compound II clearly depended on the H₂O₂ concentration, and from the biphasic time traces (Fig. 6B), k<sub>app</sub> of the dominating initial reaction was calculated to be (1.3 ± 0.03) × 10<sup>4</sup> M<sup>-1</sup>s<sup>-1</sup> (Fig. 6C).

Reaction of hsPxd01-con4-Compound I with Halides and Thiocyanate—Next, we probed the reactivity of hsPxd01-con4 compound I with the halides chloride, bromide, iodide, and the pseudo-halide thiocyanate. Again, the sequential mode was used to form compound I by preincubating hsPxd01-con4 with an equimolar concentration of hydrogen peroxide. The ferric form of hsPxd01-con4 is depicted in green, and compound II, compound III, and compound III are shown in red, blue, and cyan, respectively. The characteristic Soret peak maxima and bands in the visible region are illustrated in the same color code.

Compound II was formed by the addition of 50 µM hydrogen peroxide for 200 ms in the aging loop. The time trace of reaction between 500 nM compound I of hsPxd01-con4 and 300 µM hydrogen peroxide measured in the sequential stopped-flow mode (delay time of 200 ms for compound I formation). The time trace (black line) was fitted double exponentially (red line). C, pseudo-first-order rate constant of 500 nM hsPxd01-con4 compound I reacting with 50, 100, 150, 200, 250, and 300 µM hydrogen peroxide, respectively. k<sub>app</sub> values of the first phase were plotted against the concentration of hydrogen peroxide.

By contrast, the reaction of hsPxd01-con4 compound I with thiocyanate and iodide was very fast and resulted in a direct conversion of compound I to the ferric state. The spectral transition was almost identical to that observed with bromide. Fig. 8, A and B, shows representative monophasic time traces that could be fitted single exponentially. The apparent bimolecular rate constants were calculated to be (1.8 ± 0.07) × 10<sup>7</sup> M<sup>-1</sup>s<sup>-1</sup> and (1.7 ± 0.067) × 10<sup>7</sup> M<sup>-1</sup>s<sup>-1</sup> for the reaction with thiocyanate and iodide at pH 7.4 (Fig. 8, C and D).

**Discussion**

For the first time, a truncated variant of hsPxd01 was produced in appreciable yield, good quality, and high activity, which allowed for pre-steady-state kinetic measurements to evaluate the substrate specificity and the mechanism of halide oxidation of human peroxidasin 1. So far, only steady-state (5) and end point measurements (1, 2, 5) were published, and it was suggested that the enzyme preferentially generates hypobromous acid as a reactive intermediate to form sulfilimine cross-links in collagen (1, 2). Additionally, some papers reported the generation of hypochlorous acid by hsPxd01 (15, 16).

Human peroxidasin 1 is a homotrimeric, highly glycosylated multidomain peroxidase, which so far could only be produced in recombinant form in animal cell cultures in very low amounts of protein with unsatisfactory heme occupancy and incomplete post-translational heme modification and thus low activity (5). However, elimination of the LRR and VWC domains increased the activity of the respective recombinant
Recently, Colon and Bhave (8) demonstrated that proprotein convertase processing enhances peroxidasin 1 activity by elimination of the VWC domains and proposed that this event represents a key step in the biosynthesis and function of hsPxd01 to support basic membrane and tissue integrity. These findings motivated us to design several constructs, including the POX domain only to search for a functional and well folded protein for first comprehensive pre-steady-state kinetic studies. Finally, comparative biochemical studies demonstrated that only the construct hsPxd01-con4 fulfilled the requirements with regard to yield, heme occupancy, and modification. Apparently, the four Ig domains together with the peroxidase domain are the smallest active entity, which is also supported by data from Ero-Tolliver et al. (17) that showed that this construct is likewise the smallest unit that mediates efficient sulfilimine cross-linking. The POX domain only was inactive in these studies, which underlines the evolutionarily conserved function of peroxidasin in tissue development and integrity and distinguishes peroxidasin from other peroxidases, such as LPO, EPO, and MPO, which are composed of fully functional POX domains only.

Our spectral, redox, and kinetic data clearly demonstrate that hsPxd01-con4 has an LPO-like heme environment that was already proposed by sequence alignment and homology modeling (4, 5). It has been demonstrated that one of the most important structural features of halogenating enzymes like LPO, EPO, and MPO is the modification of the 1- and 5-methyl groups on pyrrole rings A and C of the heme group allowing formation of ester linkages with the carboxyl groups of conserved aspartate and glutamate residues (7, 18). Myeloperoxidase is unique in having a third covalent (i.e. sulfonium ion) bond (9, 19, 20). Formation of these covalent heme-protein bonds has been proposed to occur autocatalytically (22–24) mediated by (sub)micromolar hydrogen peroxide concentrations and has a deep impact on the biochemical and biophysical properties of these peroxidases (18, 19). Full establishment of the covalent bonds is never achieved even when native proteins are purified from natural sources (20, 22–24). Similarly, in the
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**TABLE 1**

| Compound         | LPO | EPO | MPO |
|------------------|-----|-----|-----|
| Fe(II)/Fe(III)   |     |     |     |
| Reduction potential | -0.128 V | -0.183 V | -0.176 V |
| Substrate        |     |     |     |
| H₂O₂             | 1800 | 1100 | 4300 |
| Cl⁻               | -   | -   | 0.31 |
| Br⁻               | 560 | 4.1 | 1900 |
| I⁻                | 1680 | 1200 | 9300 |
| SCN⁻              | 1830 | 20000 | 10000 |

Fe(III)/Fe(II) reduction potential and apparent second-order rate constants of Compound I formation (H₂O₂) and reduction (chloride, bromide, thiocyanate, and iodide) reactions of hsPxd01-con4, LPO, EPO, and MPO.

For hsPxd01-con4, all measurements were performed in 100 mM phosphate buffer, pH 7.4, whereas the data displayed for LPO, EPO, and MPO was measured in 10 mM phosphate buffer, pH 7.

In any case, we could demonstrate that ferric hsPxd01-con4 exhibits spectral features very similar to LPO and EPO and in addition shows similar rates of cyanide binding, which clearly suggests comparable active site architectures. The standard reduction potential $E^{00}$ [Fe(III)/Fe(II)] of our construct follows the hierarchy $E^{00}$ (MPO) $> 5$ mV $> E^{00}$ (hsPxd01-con4; $-128$ mV $> E^{00}$ (EPO; $-176$ mV $> E^{00}$ (LPO; $-183$ mV) (Table 1) (9, 11, 29).

The reaction cycle of hsPxd01 starts by reaction of the Fe(III) form with hydrogen peroxide to form compound I (oxoiron(IV) with porphyrin π-cation radical), which contains two oxidizing equivalents more than the resting enzyme (Reaction 1). The determined $k_{app}$ value of this bimolecular reaction was similar to that of other mammalian heme peroxidases with reported $k_{app}$ values within $(1.1–5.6) \times 10^7$ M⁻¹ s⁻¹ (14, 30, 31). Heterolytic cleavage of hydrogen peroxide is supported by a fully conserved distal His-Arg pair (His$^{287}$ and Arg$^{277}$ in hsPxd01) (30), with His$^{287}$ acting as proton acceptor and donor in this redox reaction. Upon its protonation ($pK_a \sim 4.7$) Reaction 1 cannot take place. PorFe is equal to heme or protoporphyrin IX plus iron.

\[
[\text{Por Fe(III)}] + \text{H}_2\text{O}_2 \rightarrow [\text{Por}^+ \text{Fe(IV)}=\text{O}] + \text{H}_2\text{O}
\]

**REACTION 1**

Similarly to LPO (14) and thyroid peroxidase (TPO) (32), compound I can be produced with equimolar H₂O₂. In the absence of an exogenous electron donor, it slowly converts to a compound II-like species, which most probably is compound I* formed by intramolecular electron transport from the protein matrix (where aa is amino acid) to the prosthetic group (Reaction 2).

\[
[\text{Por}^+ \text{Fe(IV)}=\text{O} \text{aa}] \rightarrow [\text{Por Fe(IV)}–\text{OH} \text{aa}^+]
\]

**REACTION 2**

Interestingly, hydrogen peroxide also mediates the one-electron reduction of compound I of hsPxd01-con4 to compound II, i.e. [Por Fe(IV)-OH] (Reaction 3).

\[
[\text{Por}^+ \text{Fe(IV)}=\text{O}] + \text{H}_2\text{O}_2 \rightarrow [\text{Por Fe(IV)}–\text{OH}] + \text{O}_2^- + \text{H}^+
\]

**REACTION 3**

which so far has been described for MPO only (30). At high (>1000) molar excess of H₂O₂, compound II is converted to compound III (Reaction 4).

\[
[\text{Por Fe(IV)}–\text{OH}] + \text{H}_2\text{O}_2 \rightarrow \text{H}_2\text{O} + [\text{Por Fe(II)}–\text{O}_3^-]
\]

\[\leftrightarrow [\text{Por Fe(III)}–\text{O}_2^-]
\]

**REACTION 4**

However, it is unlikely that this reaction is relevant in vivo because the extracellular H₂O₂ concentration is typically in the low micromolar range. Moreover, it is unknown whether there is a distinct (inducible?) source of hydrogen peroxide in the extracellular matrix for initiation of Reaction 1 and, finally, the halogenation cycle for sulfilimine formation.

It has been reported that the pseudohalide thiocyanate (SCN⁻) and iodide inhibited sulfilimine cross-linking in cell culture, whereas bromide enhanced cross-link formation (2). Now with our stopped-flow data, we can easily explain these observations. Direct reduction of compound I by halides (X⁻) or SCN⁻ restores the enzyme in its resting state and releases hypohalous acids (HOX) or hypothiocyanite (HOSCN) (Reaction 5).

\[
[\text{Por}^+ \text{Fe(IV)}=\text{O}] + \text{X}^- + \text{H}^+ \rightarrow [\text{Por Fe(III)}] + \text{HOX}
\]

**REACTION 5**
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At physiological pH 7.4 both thiocyanate (1.83 × 10^7 M⁻¹ s⁻¹) and iodide (1.68 × 10^7 M⁻¹ s⁻¹) are excellent two-electron donors of hsPxd01-con4 compound I and thus effectively compete with bromide. Reduction of compound I to the ferric resting state mediated by bromide (5.6 × 10^6 M⁻¹ s⁻¹) is about 3-times slower compared with I⁻ and SCN⁻. Importantly, chloride (even at physiological concentrations, > 100 mM) could not mediate Reaction 5. Moreover, in the presence of chloride only compound I decayed to compound I⁺ according to Reaction 2, whereas in the presence of 100 mM Cl⁻ and micromolar bromide Reaction 5 was followed. This data fit with (i) the observation that chloride did not support cross-link formation whereas addition of micromolar Br⁻ rescued sulfilimine formation (2) and with (ii) the fact that E⁰° [Fe(III)/Fe(II)] of hsPxd01-con4 is significantly less positive compared with that of MPO, which is the only known human enzyme that is able to oxidize chloride at reasonable rate at neutral pH. This enzymatic property is closely related to the MPO-specific covalent heme-protein sulfonium ion linkage which does not exist in hsPxd01 (3, 4, 9). Nevertheless, hsPxd01-con4 outperforms the reactivity of LPO and MPO toward bromide at neutral pH (Table 1).

Typical normal human serum concentrations of bromide are in the range 10–100 μM, and the Br⁻ level is maintained via diet and renal excretion (33). In vitro studies on sulfilimine formation together with modeling of the cross-linking reaction clearly demonstrated that hypobromous acid is responsible for the formation of a bromosulfonium-ion intermediate that energetically selects for sulfilimine formation (2). Based on the demonstration that (i) bromine deficiency leads to physiological dysfunction, (ii) that repletion of the element reverses dysfunction, and that (iii) biochemical data can explain the physiological function, bromine has to be considered to be an essential trace element in animals (2). Its oxidation by hsPxd01 according to Reaction 5 provides the basis for the biosynthesis of sulfilimine cross-linked collagen IV scaffolds that are central to the formation and function of basement membranes in animals (1, 2).

However, because bromine has not been considered as an essential trace element until recently, systematic investigations on its replacement have not been pursued in various disease states associated with bromide deficiency. Functional Br⁻ deficiency may occur in smokers despite normal Br⁻ levels because of elevated levels of serum SCN⁻. Normally, the level of thiocyanate in blood plasma varies in individuals from 20 to 120 μM depending on their diet but can be significantly increased in smokers (34). Under these conditions, SCN⁻ would be the preferred electron donor for compound I, and reinforcement of collagen IV scaffolds with sulfilimine cross-links may be substantially reduced. Indeed, smoking has been associated with changes in the architecture of basement membranes (35). Oxidation of SCN⁻ generates hypothiocyanate, which is a milder oxidant than hypobromous acid and reacts with thiol residues mainly (36, 37) but cannot mediate the formation of sulfilimine cross-links. Because iodide levels are below micromolar concentrations in blood plasma, I⁻ will not compete with Br⁻ for hsPxd01 compound I.

Summing up, we were able to recombinantly produce a fully functional truncated human peroxidisin 1 variant with post-translationally modified and cross-linked heme. It allowed for the first time the determination of apparent bimolecular rate constants of all relevant redox steps of the physiologically relevant halogenation cycle, i.e. the H₂O₂-mediated compound I formation followed by two-electron reduction of compound I by bromide, iodide, and thiocyanate. Besides EPO and MPO (Table 1), human peroxidisin 1 is shown to be the most effective generator of hypobromous acid in the human body.

Experimental Procedures

Materials—Bovine lactoperoxidase (L2005), sodium chloride, potassium thiocyanate, sodium cyanide, and hydrogen peroxide (30% solution) were purchased from Sigma. The concentration of hydrogen peroxide was determined at 240 nm using the molar extinction coefficient of 39.4 M⁻¹ cm⁻¹ (38). Potassium bromide and potassium iodide were obtained from Merck. All other chemicals, if not stated otherwise, were purchased from Sigma at the highest grade available. H₂O₂ solutions and potassium iodide were prepared fresh before use.

Cloning of hsPxd01-con4—Cloning, transient transfection, and expression of hsPxd01-con4 was described previously (5). The work presented here was performed with the N-terminal polyhistidine tag version of hsPxd01-con4, resulting in a translation product of 1069 amino acid residues (Pro₁²⁴₆₋Asp₁³¹₄). All amino acid residue numberings refer to the full-length hsPxd01, including the signal peptide.

Purification of hsPxd01-con4—The cell supernatant was harvested and filtrated with a 0.45-μm PVDF membrane (Duranopore) and stored at −30 °C until further processing. After thawing, the supernatant was stirred for 48 h at 4 °C before the volume was decreased (~25 times), and the cell culture medium was replaced with 100 mM phosphate buffer, pH 7.4, using a Millipore Labscale™ TFF diafiltration system. 5 ml of HisTrap™ FF columns (GE Healthcare) loaded with nickel chloride were used for the purification of hsPxd01-con4. The column was equilibrated with 100 mM phosphate buffer, pH 7.4, containing 1 M NaCl and 5 mM imidazole. The sample was adjusted to 1 M NaCl and 5 mM imidazole before loading, and the column was washed with equilibration buffer after sample loading. The protein was eluted by applying two consecutive gradients of 0–8% (2 ml/min, 10 min) and 8–70% (1 ml/min, 50 min) of 100 mM phosphate buffer, pH 7.4, containing 500 mM NaCl and 500 mM imidazole, respectively. Eluted fractions were analyzed by UV-visible spectroscopy, SDS-PAGE, and Western blotting following standard procedures (PentarHis Antibody, BSA-free from Qiagen; anti-mouse antibody, alkaline phosphatase-conjugated).

Enhanced chemiluminescence was used for the detection of covalent heme to protein linkages as described earlier (5). Fractions were pooled accordingly and concentrated in a 10-kDa molecular mass cutoff dialysis tubing (SnakeSkin™, Thermo Fisher Scientific) by applying PEG (20 kDa) to the outside of the tubing. Subsequently, the sample was dialyzed against 100 mM phosphate buffer, pH 7.4, and stored at −30 °C.

Spectral Characterization of hsPxd01-con4—The extinction coefficients of hsPxd01-con4 were determined to be 147,500
m⁻¹ cm⁻¹ at 280 nm and 101,400 m⁻¹ cm⁻¹ at the heme Soret peak, resulting in a theoretical purity number of 0.7 (ε₄₁₂ nm/ε₂₈₀ nm). The average purity number obtained by metal affinity chromatography was 0.45–0.55 indicating a 65–80% heme occupancy. Specified hsPxd01-con4 concentrations were always related to heme concentrations.

Spectroelectrochemistry—The standard reduction potential (E°) of the Fe(III)/Fe(II) couple of hsPxd01-con4 was determined as described previously (5). Briefly, the spectroelectrochemical titrations were performed using a homemade OTTLE (optically transparent thin layer spectroelectrochemical) cell. The three-electrode configuration consisted of a gold mini-grid working electrode (Buckbee-Mears), a saturated calomel (Hg₂Cl₂) microreference electrode (AMEL Electrochemistry), separated from the working solution by a Vycor set, and a platinum wire as counter-electrode (11, 21, 29). All potentials are referenced to the standard hydrogen electrode.

Experiments were performed with 5 μM hsPxd01-con4 in 100 mM phosphate buffer, pH 7.4, containing 100 mM NaCl, 30 mM methyl viologen, and 1 μM lumiflavin 3-acetate, methylene blue, phenazine methosulfate, and indigo disulfonate used as mediators at 25 °C. Nernst plots consisted of at least five points and were invariably linear with a slope consistent with the expected slope of 6 mV.

Stopped-flow Spectroscopy—Pre-steady-state spectra were recorded with the stopped-flow apparatus SX.18MV (Applied Photophysics) connected to a diode array detector (DAD) with the first spectrum usually recorded 3 ms after mixing the reactants. The Pi-star-180 apparatus from Applied Photophysics was employed for all single wavelength measurements, and the first data point after mixing two solutions was typically recorded at 1 ms. The optical quartz cell had a volume of 20 μl and a path length of 10 mm. All reactions were followed at single wavelengths and additionally by using the DAD. Polychromatic data were analyzed with the Pro-Kineticist software from Applied Photophysics. Rate constants were determined by fitting single wavelength time traces with the Pro-Data Viewer software (Applied Photophysics). The conventional mode was applied to monitor the reaction of hsPxd01-con4 with hydrogen peroxide by following the decrease of absorbance at 412 nm and cyanide binding by monitoring the increase at 434 nm. All presented rate constants were measured using the sequential mixing mode due to the inherent instability of compound I. A delay time of 200 ms for the formation of compound I was employed.

All reactions with the exception of the pH profiles presented were performed in 100 mM phosphate buffer, pH 7.4, and at 25 °C. Citrate phosphate buffer was used for measurements from pH 4 to 5.5; phosphate buffer was employed for the pH range of 5.5–8, and carbonate buffer was used for pH 9 and 10. Three measurements were performed for each ligand (cyanide), oxidant (hydrogen peroxide), and electron donor (halides and thiocyanate) concentration, respectively. The mean of the first-order rate constants, kₐₑₛₜ, was used to calculate the apparent second-order rate constant that was obtained from the slope of the plot of the kₑₛₜ values versus the concentrations of the respective reactants.

Author Contributions—P. G. F. and C. O. conceived and coordinated the study and wrote the paper. M. P. P. designed the constructs, performed and analyzed the experiments, and contributed to the writing of the paper. R. S. K., I. S., E. E., B. S., and M. S. provided technical assistance and produced and purified the recombinant proteins. M. B. and G. B. performed the spectroelectrochemical experiments, and S. H. probed the homogeneity and conformational stability of the constructs.

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