A Chemically Programmed Proximal Ligand Enhances the Catalytic Properties of a Heme Enzyme

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ABSTRACT: Enzymes rely on complex interactions between precisely positioned active site residues as a mechanism to compensate for the limited functionality contained within the genetic code. Heme enzymes provide a striking example of this complexity, whereby the electronic properties of reactive ferryl intermediates are finely tuned through hydrogen bonding interactions between proximal ligands and neighboring amino acids. Here, we show that introduction of a chemically programmed proximal N\textsubscript{ω}-methyl histidine (NMH) ligand into an engineered ascorbate peroxidase (APX2) overcomes the reliance on the conserved Asp-His hydrogen bonding interaction, leading to a catalytically modified enzyme (APX2 NMH), which is able to achieve a significantly higher number of turnovers compared with APX2 without compromising catalytic efficiency. Structural, spectroscopic and kinetic characterization of APX2 NMH and several active site variants provides valuable insights into the role of the Asp-His-Fe triad of heme peroxidases. More significantly, simplification of catalytic mechanisms through the incorporation of chemically optimized ligands may facilitate efforts to create and evolve new active site heme environments within proteins.

INTRODUCTION

Enzymes are able to mediate a diverse array of energetically demanding transformations using a limited set of functional amino acids presented by the genetic code. The enormous rate enhancements provided by these biological catalysts are achieved through the evolution of sophisticated catalytic machineries that typically rely on interactions between precisely positioned networks of amino acids. Considerable effort has been devoted to understanding the catalytic roles played by the individual functional amino acids across the different enzyme classes. Perhaps not surprisingly, such analysis reveals that histidine is the most frequently encountered and versatile histidine provides an essential role in catalysis. For example, the conserved Asp-His-Ser triad of serine proteases and the Asp-His-Fe triad of heme peroxidases are among the most studied catalytic motifs in biochemistry (Figure 1a). While the primary function of the central histidine residue in such systems is often well established, defining the precise role played by secondary interactions with neighboring residues during catalysis presents a formidable challenge.

Mutation of an amino acid that is intimately involved in an enzyme’s catalytic mechanism typically results in dramatic reduction in activity, providing a simple method of identifying essential residues. However, the limited range of functionality present within the canonical amino acids severely restricts the degree of mechanistic information that can be uncovered using this approach. In the present study, we demonstrate how expansion of the genetic code to include structural analogues of histidine provides an effective strategy to probe key secondary interactions required for efficient catalysis in a heme enzyme. Such fundamental understanding is essential to advance our ability to rationally re-engineer existing enzymes or design biocatalysts with novel function.

Heme peroxidases utilize hydrogen peroxide as an electron acceptor to catalyze a range of oxidative transformations that have been exploited for biotechnological applications. These enzymes utilize a histidine residue as the axial ligand of heme iron, with a hydrogen bond formed between the non-coordinating N\textsubscript{ω} atom and an aspartate residue. This Asp-His-Fe triad is conserved across the heme peroxidase family

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Despite the similar active site architectures of CCP and APX, the electronic properties of the compound I intermediate of these two enzymes differ significantly, with a proximal Trp191 centered radical cation formed in CCP compared with a porphyrin π cation radical in APX.19–21 In short, the complex network of interactions formed by the proximal aspartate residue make it challenging to define the importance of the His-Asp hydrogen bond by simple mutation of this conserved amino acid.

Expansion of the genetic code to include a range of histidine analogues has recently been described,22 presenting an opportunity to study the significance of the Asp-His hydrogen-bonding interaction from a different angle, namely by direct replacement of the axial histidine ligand with structurally related analogues. A number of features make N\textsubscript{3}-methyl histidine (NMH) a particularly attractive residue to probe Asp-His interactions. First, the absence of the N\textsubscript{3} hydrogen means that NMH is incapable of forming a hydrogen bond to the aspartate residue. Due to the comparable pK\textsubscript{a} values associated with imidazole and N\textsubscript{3}-methylimidazole (7.18 and 7.38, respectively),23 relatively minor differences in electron donating properties of proximal histidine and NMH ligands are expected in the absence of additional interactions. Finally, the alkyl modification effectively forces the imidazole ring to adopt a single tautomeric form and prevents the formation of “imidazole like” proximal ligands (via loss or polarization of the N\textsubscript{3} hydrogen), thus overcoming ambiguity surrounding imidazole charge state which complicates interpretation of catalytic mechanisms involving histidine.14 Here, we demonstrate that substitution of the proximal histidine with NMH in an engineered ascorbate peroxidase (APX2) results in a modified enzyme with enhanced catalytic properties, despite disrupting the “essential” Asp-His hydrogen bonding interaction which is conserved across the peroxidase family.

### RESULTS

An engineered variant of ascorbate peroxidase24 was selected as a model enzyme for proximal ligand replacement (Figure 2a). The selective incorporation of NMH into proteins in response to amber codons was recently described using an evolved pyrrolysyl-tRNA synthetase/pyrrolysl-tRNA (PylRS(NMH)/tRNA\textsubscript{Pyl}) pair.22 Two copies of the relevant PylRS variant were subcloned into the pEVL plasmid which has been optimized for the production of proteins containing non-canonical residues.25 Coexpression of this pEVL construct with APX2 (AMBER) on a standard pET29b vector led to the production of the desired APX2 variant, with a protein yield of >50 mg per liter of culture after purification (see Experimental Procedures). Stoichiometric replacement of the proximal histidine residue (His163) with NMH was confirmed by MS analysis of the intact protein as well as peptide fragments generated by tryptic digestion (Figure S1). The UV–vis spectrum of His163NMH APX2 (APX2 NMH) in the resting state is indistinguishable from that of the wild-type enzyme, suggesting that this proximal ligand modification causes minimal changes to the electronic and structural properties of the heme iron (Figure 2b). Both variants are characterized by a Soret maximum at 404 nm with associated Q bands at 500/632 nm and 540/574 nm, consistent with the presence of a mixture of six-coordinated (6c) high-spin (HS) and low-spin (LS) heme iron.26 The heme occupancy of purified APX2 and APX2 NMH variants is also very similar, as judged by the ratio of intensities of the bands at 404 and 280 nm. Addition of cyanide...
to the resting enzymes results in similar spectral changes in APX2 and APX2 NMH due to the formation of 6c-LS species (Figure 2c). Compounds I and II (Figure 1b) have been implicated as common intermediates in many proteins. Although these species are highly unstable in enzymes like the cytochrome P450s, they are readily detected in APX under relatively mild conditions. The UV−vis spectra of compound I (Figure 2d) and compound II (Figure 2e) intermediates of APX2 and APX2 NMH were therefore recorded following mixing of the resting enzymes with hydrogen peroxide in the absence of phenolic substrate. Only minor differences are observed between the spectral features of these key APX2 and APX2 NMH ferryl intermediates, suggesting that the proximal His → NMH substitution does not lead to significant changes in the electronic structures of compound I and compound II.

The catalytic parameters of the APX2 and APX2 NMH variants for the oxidation of the peroxidase substrate o-methoxyphenol (guaiacol) in the presence of hydrogen peroxide (2.5 mM) were determined using a stopped-flow apparatus. Peroxidase-mediated guaiacol oxidation leads to the formation of a stable tetrameric species whose formation can be readily monitored by absorbance at 470 nm (Scheme 1). This assay has been exploited previously for the characterization of APX2, with an optimal H2O2 concentration between 1.0 and 10 mM reported. Remarkably, despite disruption of the conserved Asp-His hydrogen bonding interaction, introduction of a proximal NMH ligand results in a slight increase in catalytic efficiency, with $k_{cat}/K_m$ values of $8.6 \times 10^4 \text{M}^{-1} \text{s}^{-1}$ ($K_m = 36 \pm 5 \text{mM}$ and $k_{cat} = 3060 \pm 360 \text{s}^{-1}$) and $1.1 \times 10^5 \text{M}^{-1} \text{s}^{-1}$ ($K_m = 14 \pm 1 \text{mM}$ and $k_{cat} = 1500 \pm 90 \text{s}^{-1}$) determined for APX2 and APX2 NMH, respectively (Figure 3a). Furthermore, in the absence of substrate under presteady-state conditions, only minor differences in the rate constants for compound I ($k_1$) and compound II ($k_2$) formation were determined for APX2 ($k_1 = 1.4 \times 10^9 \text{M}^{-1} \text{s}^{-1}$, $k_2 = 1.2 \times 10^8 \text{M}^{-1} \text{s}^{-1}$) and APX2 NMH ($k_1 = 6.0 \times 10^8 \text{M}^{-1} \text{s}^{-1}$, $k_2 = 6.1 \times 10^7 \text{M}^{-1} \text{s}^{-1}$) (Figure S2).

Determination of the total turnovers achieved by APX2 and APX2 NMH reveals a more dramatic difference in their catalytic behavior. Heme peroxidases are known to suffer irreversible inactivation during catalysis (phenol oxidation), a significant drawback when potential industrial applications are considered. In our hands, APX2 was able to promote guaiacol oxidation with a total turnover number (TON) of 6200 prior to irreversible inactivation, a value comparable to that reported previously, albeit under slightly modified conditions (Figure 3b). In contrast, a TON of 31 300 was determined for APX2 NMH, which represents a 5-fold increase as a result of the His → NMH proximal ligand mutation (Figure 3b). The UV−vis
Figure 4. Crystal structures of APX and APX2 NMH. (a) The active site of APX (PDB ID: 1OAG). The heme cofactor and the proximal His163, Trp179 and Asp208 are shown in atom colored sticks (carbon, nitrogen and oxygen colored in magenta, blue and red, respectively) with the remainder of the protein showed in ribbon (colored cyan). An ordered water molecule is present in the distal pocket binding the heme iron (shown as a red sphere). (b) The active site of APX2 NMH. The heme cofactor and the proximal NMH163, Trp179 and Asp208 are shown in atom colored sticks (carbon, nitrogen and oxygen colored in green, blue and red, respectively) with the remainder of the protein showed in ribbon (colored in brown). The omit maps for the heme cofactor, NMH163, and Asp208 shown as gray mesh, contoured at 2.5 s.u. An ordered water molecule is present in the distal pocket binding the heme iron (shown as a red sphere). (c) Overlay of the active sites of APX and APX2 NMH to minimize the rmsd between the heme cofactors. The heme cofactors, active site residues and ordered water molecules of APX and APX2 NMH are shown in atom colored sticks, color coded as in panels a and b, respectively.

Table 1. Catalytic Parameters for Asp208 Variants of APX2 and APX2 NMH

| variant       | \( k_{cat} \) (s\(^{-1}\)) | \( K_{m} \) (mM) | \( V_{max} \) (mM) | \( k_{cat}/K_{m} \) (mM\(^{-1}\) s\(^{-1}\)) | total TON |
|---------------|-----------------------------|-----------------|-----------------|----------------------------------|---------|
| APX2          | 3060                        | 36.0            | 12.5            | 86                               | 6200    |
| APX2 D208N    | 8.8                         | 1.5             | 25              | 6.0                              | 600     |
| APX2 D208M    | 88.7                        | 19.0            | 103             | 4.7                              | 1000    |
| APX2 NMH      | 1500                        | 14.0            | 14.1            | 110                              | 31 300  |
| APX2 NMH D208A| 194                         | 11.0            | 74.8            | 18                               | 7000    |
| APX2 NMH D208N| 351                         | 18.5            | 44.9            | 19                               | 9600    |
| APX2 NMH D208M| 780                         | 25.9            | 34.1            | 30                               | 18 900  |

All measurements were carried out at 25 °C in PBS containing 2.5 mM H\(_2\)O\(_2\) and 5 nM enzyme. \( k_{cat}/K_{m} \) was derived from the Michaelis–Menten equation for uncompetitive substrate inhibition.
variants, and significantly higher than the corresponding APX2 D208M variant ($k_{cat}/K_m = 4.7 \times 10^3 \text{ M}^{-1} \text{s}^{-1}$; TON; 1000) (Figure 5 and S3). Interestingly, a methionine residue has been identified at the corresponding position of dehaloperoxidase hemoglobin A (DHP A). In this enzyme, one of the sulfur lone pairs is in contact with the proximal histidine ligand which has been proposed to increase the electron donating character of the histidine through a charge transfer interaction. It is conceivable that the methionine residue performs a similar role in the APX2 NMH D208M variant. Combined, these data suggest that replacement of the axial histidine ligand with NMH confers a greater degree of mutational flexibility at position 208. Despite the fact that it is no longer able to participate in a hydrogen-bonding interaction with the proximal ligand, aspartate still remains the optimal residue at position 208. We speculate that this residue may stabilize partial positive charge that develops on the axial histidine ligand through electrostatic interactions or, alternatively, play a role in optimizing the $\pi-\pi$ interaction between Trp179 and the proximal ligand.

In contrast to APX2 and APX2 NMH which display indistinguishable spectral features, comparison of the UV–vis spectra of the D208X variants reveal clear differences (Figure S5). Mutation of Asp208 in APX2 NMH causes displacement of the Soret maximum from 404 to 408 nm. The four Q bands (at 502/633 nm and 540/576 nm) associated with 6c-HS and 6c-LS heme iron, respectively, are evident in all D208X variants investigated (D208M, D208N and D208A), albeit with a reduced 6c-LS:6c-LS ratio compared with the starting Asp208 variant. In contrast, introduction of the corresponding point mutations into APX2 (D208N and D208M) causes displacement of the Soret to 411 nm, loss of two Q bands associated with the 6c-HS species, and the emergence of a new band at 669 nm which is also present to a lesser extent in the UV–vis spectra of selected APX2 NMH D208X variants. These data suggest that the Asp208 residue may play a key role in stabilizing the catalytically productive electronic structure of the heme iron in APX2 which is less critical in APX2 NMH.

**DISCUSSION**

Considerable effort has been devoted to unraveling the precise mechanistic role played by conserved Asp-His hydrogen bonding interactions which are prevalent catalytic motifs found in diverse classes of metallo- and non-metalloenzymes. Suggested roles include controlling the conformation of the histidine side chain, stabilizing the appropriate histidine tautomer, stabilizing positively charged histidine intermediates formed along the reaction coordinate during catalysis, perturbing the $pK_a$ of the histidine residue, or a combination of the above. Specifically in the case of heme peroxidases, the Asp-His interaction is reported to impart partial imidazolate character onto the axial ligand which may assist in O–O bond heterolysis or perturb the electronic properties of the iron center.

In this study, disruption of the Asp-His hydrogen bond by replacement of the proximal histidine ligand with the noncanonical structural analogue NMH has allowed us to re-examine the mechanistic role of this conserved interaction in heme peroxidases. Replacement of the $N_2$ hydrogen of histidine with an alkyl residue has a number of structural/functional consequences which directly relate to previously suggested roles for Asp-His interactions. This modification (1) effectively fixes the tautomeric form of the imidazole ring; (2) ensures the neutral charge state of the proximal ligand throughout the catalytic cycle (i.e., formation of the anionic ligand via deprotonation of the $N_2$ hydrogen is not possible); and (3) prevents buildup of imidazolate character through $N$–H bond polarization, thus overcoming ambiguity surrounding imidazolate charge state. The comparable resting state UV–vis spectra and rates of compound I formation in APX2 and APX2 NMH suggest that any contribution made by the proximal Asp-His hydrogen bond (in APX2) to $H_2O_2$ heterolysis or to the electronic properties of the heme cofactor (resting state) is compensated by the minor increase in basicity associated with the NMH ligand. It can be assumed that the tautomeric form of the APX2 histidine ligand is controlled by coordination to the $Fe$ center (assuming that the histidine ligand is in the neutral form). Furthermore, comparison of the X-ray structure of APX2 NMH and previously reported APX structures reveals only minor structural changes which occur to accommodate the methyl substituent. The orientation of the histidine and NMH ligands is almost identical in these structures, suggesting that the Asp-His interaction is not required to control the conformation of the histidine side chain. Thus, comparison of the kinetic parameters, spectroscopic properties and X-ray structures has allowed us to exclude a number of plausible mechanistic roles for the Asp-His hydrogen-bonding interaction.

So how can we explain the increased “robustness” of APX2 NMH toward irreversible inactivation compared with APX2, and how does this relate to the role of the proximal pocket Asp-His hydrogen-bonding interaction? Several hypotheses for peroxidase deactivation mechanisms have been proposed.36,37 Our study shows that small structural/electronic changes in the proximal pocket can profoundly influence these processes. In agreement with the aforementioned theoretical studies,41–48 we suggest that the Asp-His interaction may play a role in stabilizing the reactive ferryl intermediates, potentially by promoting partial delocalization of the porphyrin radical of
compound I onto the proximal histidine ligand. In that case, the introduction of a proximal NMH ligand (in APX2 NMH) would provide a more effective and robust means of stabilizing compound I that is not reliant on a noncovalent hydrogen-bonding interaction which could potentially be susceptible to uncoupling throughout repeated catalytic cycles. The electron-donating methyl substituent on the noncoordinating N4 atom is expected to increase radical stability through hyperconjugation, thus favoring radical delocalization onto the proximal NMH residue. This mechanistic interpretation is consistent with the observation that APX2 NMH is tolerant to mutations at Asp208, while such modifications in APX2 lead to dramatic reductions in catalytic activity and total turnover number.

What is evident from the present study is that the direct targeting of “essential” catalytic residues for substitution by chemically programmed amino acids provides a fruitful avenue to probe enzyme mechanism \(^{30−42}\) and to produce biocatalysts with enhanced properties. This is in contrast to traditional enzyme engineering approaches, where canonical substitutions of catalytic amino acids often lead to a dramatic loss of enzyme activity. In this study, disruption of the catalytic Asp-His-Fe triad of a heme peroxidase by introduction of a proximal NMH ligand led to biocatalysts with comparable catalytic efficiencies to the “wild type” enzyme but with increased robustness toward irreversible inactivation during catalysis (i.e., increase in total turnover numbers). Natural evolutionary processes have clearly not optimized the proximal pocket to support this chemically modified residue, and it would be interesting to observe whether laboratory evolution can further enhance the catalytic parameters of the modified enzyme. Perhaps equally significant is the increased mutational tolerance at Asp208 as a result of introducing a chemically modified residue. This finding demonstrates how nature relies upon the evolution of complex interactions between catalytic residues as a mechanism to compensate for the limited functionality contained within the genetically encoded amino acids. Simplification of catalytic mechanisms through the genetic incorporation of chemically programmed amino acids \(^{30−42}\) will certainly enhance our understanding of the minimum requirements for catalyst design and should facilitate efforts to create and evolve new active site chemistries/environments within proteins. \(^{43}\)

### EXPERIMENTAL PROCEDURES

#### Materials

All chemical and biological reagents were purchased from commercial suppliers and were used without further purification. LB medium and 2xYT were purchased from MP Biomedicals, LLC (California, USA) and Sigma-Aldrich (Missouri, USA). Hemin chloride and 30% hydrogen peroxide were purchased from Merck and should facilitate the targeting of reductions in catalytic activity and total turnover number. Observation that APX2 NMH is tolerant to mutations at Asp208, while such modifications in APX2 lead to dramatic reductions in catalytic activity and total turnover number.

What is evident from the present study is that the direct targeting of “essential” catalytic residues for substitution by chemically programmed amino acids provides a fruitful avenue to probe enzyme mechanism \(^{30−42}\) and to produce biocatalysts with enhanced properties. This is in contrast to traditional enzyme engineering approaches, where canonical substitutions of catalytic amino acids often lead to a dramatic loss of enzyme activity. In this study, disruption of the catalytic Asp-His-Fe triad of a heme peroxidase by introduction of a proximal NMH ligand led to biocatalysts with comparable catalytic efficiencies to the “wild type” enzyme but with increased robustness toward irreversible inactivation during catalysis (i.e., increase in total turnover numbers). Natural evolutionary processes have clearly not optimized the proximal pocket to support this chemically modified residue, and it would be interesting to observe whether laboratory evolution can further enhance the catalytic parameters of the modified enzyme. Perhaps equally significant is the increased mutational tolerance at Asp208 as a result of introducing a chemically modified residue. This finding demonstrates how nature relies upon the evolution of complex interactions between catalytic residues as a mechanism to compensate for the limited functionality contained within the genetically encoded amino acids. Simplification of catalytic mechanisms through the genetic incorporation of chemically programmed amino acids \(^{30−42}\) will certainly enhance our understanding of the minimum requirements for catalyst design and should facilitate efforts to create and evolve new active site chemistries/environments within proteins. \(^{43}\)

#### Physical and Biophysical Measurements

UV–vis absorption spectra were acquired on a Lambda 35 spectrophotometer (PerkinElmer, Massachusetts, USA). Fast protein liquid chromatography (FPLC) was carried out on an NGC Quest 10 plus chromatography system (Biorad laboratories, California, USA) equipped with a multichannel detector and BioFrac fraction collector. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was performed on a PhastSystem electrophoresis apparatus using precast PhastGel homogeneous 12.5 (GE healthcare Life Sciences, Chicago, USA). Matrix-assisted laser ionization time-of-flight mass spectrometry (MALDI-TOF-MS) was performed on a microflex (Bruker Daltonics, Bremen, Germany) using 2,5-dihydroxybenzoic acid (HCA) or sinapinic acid (SA) as a matrix. High resolution mass spectra were recorded on an electrospray ionization quadrupole time-of-flight (ESI-Q-TOF) system MaXis (Bruker Daltonics, Bremen, Germany) coupled with an Agilent 1200 system (Agilent technology, USA). Steady-state and presteady-state stopped-flow experiments were performed on an Applied Photophysics SX18 stopped-flow spectrophotometer (Applied Photophysics Ltd., Leatherhead, UK) equipped with a xenon arc lamp and a 1 cm path length.

#### Construction of pET29_APX2, APX2 H163TAG and D208X Variants

The gene encoding APX2 was amplified from pCDNA3 APX2-NESt (Addgene plasmid #49386) \(^{24}\) using the primer pair APX2_NdeI_for and APX2_XhoI_rev (Table S1). The resulting DNA fragment was digested by NdeI and XhoI and ligated into a pET29b(+) vector digested by the same restriction enzymes, yielding pET29b_APX2. The codon for His163 in pET29b_APX2 was replaced by an amber codon (TAG) by overlap extension PCR. Briefly, two fragments corresponding to the N and C termini were synthesized with the primer pairs APX2_NdeI_for and H163TAG_rev, H163TAG_for and APX2_XhoI_rev (Table S1). The two PCR products were purified and then combined in an equimolar ratio for final PCR. The full length fragment was assembled using the primer pair APX2_NdeI_for and APX2_XhoI_rev. The mutated fragment was digested by NdeI and XhoI and ligated into a pET29b(+) as described above, yielding pET29b_APX2 H163TAG. The Asp208 mutations were introduced into the pET29b_APX2 and pET29b_APX2 H163TAG by overlap extension PCR. Two fragments corresponding to the N and C termini were synthesized with the primer pairs APX2_NdeI_for and D208_rev, D208A_for, D208N_for or D208M_for and APX2_XhoI_rev. The N and C terminal PCR products were purified and then combined in an equimolar ratio for final PCR. The full length fragments were assembled using the primer pairs APX2_NdeI_for and APX2_XhoI_rev. The mutated fragment was digested by NdeI and XhoI and ligated into a pET29b(+) as described above. The DNA sequence was confirmed by DNA sequencing service provided by Microsynth AG.

#### Construction of pEVOL_PylRS(NMH)

Five mutations (L207I, Y217F, L274G, C313F, and Y349F) were introduced into the pBK-PylRS plasmid generate pBK-PylRS(NMH) as described previously. \(^{22}\) The Y349F point mutation was first introduced by overlap extension PCR. Fragments corresponding to the N and C termini were synthesized with the primer pairs PylRS_for and PylRS_rev (Table S1). The products of the two previous PCRs were purified and then combined in equimolar amounts for the final PCR. The full-length fragment was amplified using the primer pair PylRS_for and PylRS_rev (Table S1). The mutated fragment was digested by NdeI and PstI and ligated into a pBK vector digested by the same restriction enzymes, yielding pBK_PylRS Y349. Three fragments corresponding to the N terminus, middle, and C terminus were synthesized with PylRS_for and PylRS_rev (Table S1). The three PCR products were purified and then combined in equimolar ratios for the final PCR. The full length fragment was assembled using a primer pair PylRS_for and PylRS_rev (Table S1). The mutated fragment was digested by NdeI and PstI and ligated into a pBK vector digested by the same restriction enzymes, yielding pBK_PylRS(NMH). To construct pEVOL_PylRS(NMH), the same mutated fragment digested by NdeI and PstI was initially ligated into a pEVOL vector digested by the same restriction enzymes. Next, the PylRS(NMH) gene was amplified using the primer pair pBK_PylRS(NMH)_BglII and pBK_PylRS(NMH)_SalI, digested with BglII and SalI, and ligated into the pEVOL vector digested by the same restriction enzymes to give pEVOL_PylRS(NMH) containing two copies of the PylRS(NMH).
gene. The DNA sequence was confirmed as described above using pBAD_for, pBAD_rev, and two custom primers (pEVOLconst_for and pEVOLconst_rev).

**Protein Production and Purification.** For expression of APX2, pET29b APX2 was transformed into BL21(DE3) E. coli and the cells were plated on a LB agar plate containing 50 μg/mL kanamycin. A single colony of freshly transformed cells was cultured overnight in 3 mL of LB medium containing 50 μg/mL kanamycin. One mL of the culture was used to inoculate 100 mL of 2xYT medium supplemented with 1 mM δ-aminolevulinic acid and 50 μg/mL kanamycin. The culture was incubated for ∼2 h at 37 °C with shaking speed at 230 rpm. When the OD600 of the culture reached ∼0.5, IPTG was added to a final concentration of 0.1 mM to induce overexpression of the APX2 gene. APX2 D208X variants were produced using this same procedure. For expression of APX2 NMH, pET29b APX2 H163TAG and pEVOL PyR3(NMH) were transformed into BL21(DE3) E. coli and the cells were plated on a LB agar plate containing 50 μg/mL kanamycin and 34 μg/mL chloramphenicol. A single colony of freshly transformed cells was cultured overnight in 3 mL of LB medium containing 50 μg/mL kanamycin and 34 μg/mL chloramphenicol. One mL of the culture was used to inoculate 100 mL of 2xYT medium supplemented with 1 mM δ-aminolevulinic acid, 12 mM NaCl, 50 μg/mL kanamycin and 34 μg/mL chloramphenicol. The culture was incubated for ∼2.5 h at 37 °C with shaking at 230 rpm. When the OD600 of the culture reached ∼0.5, IPTG and arabinose were added to a final concentration of 0.1 mM and 0.05%, respectively. APX2 NMH D208X variants were produced using this same procedure. The induced cultures were incubated for ∼24 h at 25 °C, and the cells were subsequently harvested by centrifugation at 5500g for 15 min. The pelleted bacterial cells were suspended in 50 mM Tris·HCl, 300 mM NaCl at pH8.0 (buffer A) with 10 mM imidazole containing 0.5% lysosome and 0.01% DNase and sonicated to disrupt the cells. The lysate was centrifuged at 11 500g for 15 min and the supernatant was subjected to affinity chromatography using Ni-NTA Agarose (Qiagen, Hilden, Germany). His-tagged APX2 and APX2 NMH were eluted by buffer A containing 300 mM imidazole. To maximize heme occupancy, partially purified APX2 and APX2 NMH were reconstituted with heme chloride. Briefly, Ni-NTA purified protein solutions were rapidly mixed with one equivalent of heme chloride dissolved in 10 mM NaOH, and incubated at room temperature for 1 h. The APX2 and APX2 NMH solutions were then subjected to size-exclusion (SEC) chromatography on Superdex 200 10/300 GL (GE healthcare Life Sciences) with a flow rate of 0.5 mL/min using an NGC Quest 10 plus FPLC system equilibrated with PBS (10 mM Na2HPO4, 1.8 mM KH2PO4, 2.7 mM KCl, and 137 mM NaCl, pH 7.4) at a final concentration of 2–5 μM. The CN-bound APX2 and APX2 NMH were obtained following addition of sodium cyanide at the final concentration of 1 mM in PBS (Figure 2c). Compound I and II of APX2 and APX2 NMH (Figure 2d, e) were generated as previously described. 28 Briefly, APX2 and APX2 NMH were diluted to 10 μM on the basis of the heme absorption (ε405 = 128 000 M−1 cm−1) in PBS. PBS buffer containing 20 or 500 μM H2O2 was prepared and used immediately. Solutions were loaded into the stopped-flow UV–vis spectrometer equilibrated to 25 °C. Compound I formation was complete in ∼50 ms for a 1:2 ratio of enzyme and hydrogen peroxide. Compound II formation was complete in ∼400 ms for a 1:5 ratio of enzyme and hydrogen peroxide.

**Determination of Total Turnover Numbers.** Reactions were carried out at 25 °C in PBS in 1 mL quartz cuvettes containing 10 mM guaiacol and 2.5 mM H2O2. Reactions were initiated by adding freshly purified APX2 and APX2 NMH at the final concentration of 5 nM. Product formation was monitored by the change in absorbance at 450 nm (ε450 = 8209 M−1 cm−1).

**Stopped-Flow Kinetics.** APX2, APX2 NMH and D208X variants were diluted to 10 nM on the basis of the heme absorption (ε405 = 128 000 M−1 cm−1) in PBS. Substrate solutions containing 1, 2.5, 5, 7.5, 10, 15, 20, 30, 40, 60, and 100 mM guaiacol and 5 mM H2O2 were prepared in PBS and used immediately. Solutions were loaded into the stopped-flow UV–vis spectrometer equilibrated to 25 °C. The product formation was monitored by the change in absorbance at 470 nm. Typically, 3–5 shots were averaged and the averaged traces were fitted by linear regression using Kaleidograph. The observed initial rates were fitted to the Michaelis–Menten equation for uncompetitive substrate inhibition (Figure 3a for APX2 and APX2 NMH; Figure S3 for APX2 NMH D208X, D208N, D208A, and APX2 D208M, D208N). Pseudo-first-order rate constants for compound I formation in APX2 and APX2 NMH were obtained as follows. Briefly, one syringe of the stopped-flow apparatus containing 2 μM enzyme in PBS was mixed with the other containing at least a 10-fold excess of H2O2 (20, 30, 40, and 50 μM) in the same buffer. The second order rate constants were extracted from the slope of the plots. 28 Compound I formation was monitored by a decrease in absorbance at 398 nm. Typically, compound II formation was monitored by an increase in absorbance at 424 nm following mixing of 4 μM enzyme in PBS with an equal volume of 300 μM H2O2 in the same buffer.

**Substrate Scope of APX2 and APX2 NMH.** Reactions were carried out at 25 °C in PBS in 1 mL quartz cuvettes containing 10 mM substrates and 2.5 mM H2O2. Reactions were initiated by adding freshly purified APX2 and APX2 NMH at the final concentration of 5 nM. Product formation was monitored by the change in absorbance at 480, 410, and 420 nm for o-anisidine, o- cresol, and 3,5-dimethylanisole, respectively. The relative number of turnovers performed by APX2 NMH and APX2 are shown in Figure 3c. For the experiments with phenol, a stopped-flow UV–vis spectrometer was used due to the relatively rapid decay of the products formed. Briefly, APX2 and APX2 were diluted to 10 nM on the basis of the heme absorption in PBS. Substrate solutions containing 20 mM phenol and 5 mM H2O2 were prepared in PBS and used immediately. Solutions were loaded into the stopped-flow UV–vis spectrometer equilibrated to 25 °C. Product formation was monitored by the change in absorbance at 420 nm. The relative number of turnovers performed by APX2 NMH and APX2 are shown in Figure 3c.

**Crytalization and Structure Determination.** APX2 NMH in PBS buffer was concentrated to approximately 20 mg/mL in an
Amicon Ultra-15 10-kDa cutoff centrifugal filter device. The protein solution was passed through a PD-10 column (GE Healthcare Life Sciences) pre-equilibrated with pure water, and the final concentration was adjusted to 15 mg/mL. The protein was crystallized by vapor diffusion in sitting drops at 20 °C. An initial crystallization hit was identified in a well containing 100 mM Bis-Tris at pH 5.5 and 2.0 M ammonium sulfate using a commercial sparse matrix screen (JCSG+ Suite, Qiagen). Crystallization conditions were improved by varying the buffer pH and ammonium sulfate concentration. The optimal crystallization condition was 100 mM bis-tris at pH 7.0 containing 2.0 mM ammonium sulfate. Crystals were cryoprotected by plunging into mother liquor supplemented with 20% w/v glycerol prior to flash freezing in liquid nitrogen. Diffraction data were collected at 100 K, corrected for Lorentz and polarization effects, and then indexed and integrated using MOSFLM. The space group was determined to be P2₁2₁2₁ by the change in absorbance at 470 nm (Figure S4).

The crystal structure of APX2 NMH was deposited in the RCSB Protein Data Bank (PDB) under accession number 5L8E.

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