A Noncanonical Tryptophan Analogue Reveals an Active Site Hydrogen Bond Controlling Ferryl Reactivity in a Heme Peroxidase


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ABSTRACT: Nature employs high-energy metal-oxo intermediates embedded within enzyme active sites to perform challenging oxidative transformations with remarkable selectivity. Understanding how different local metal-oxo coordination environments control intermediate reactivity and catalytic function is a longstanding objective. However, conducting structure–activity relationships directly in active sites has proven challenging due to the limited range of amino acid substitutions achievable within the constraints of the genetic code. Here, we use an expanded genetic code to examine the impact of hydrogen bonding interactions on ferryl heme structure and reactivity, by replacing the N−H group of the active site Trp51 of cytochrome c peroxidase by an S atom. Removal of a single hydrogen bond stabilizes the porphyrin π-cation radical state of CpW191F compound I. In contrast, this modification leads to more basic and reactive neutral ferryl heme states, as found in CpW191F compound II and the wild-type ferryl heme-Trp191 radical pair of compound I. This increased reactivity manifests in a >60-fold activity increase toward phenolic substrates but remarkably has negligible effects on oxidation of the biological redox partner cyt c. Our data highlight how Trp51 tunes the lifetimes of key ferryl intermediates and works in synergy with the redox active Trp191 and a well-defined substrate binding site to regulate catalytic function. More broadly, this work shows how noncanonical substitutions can advance our understanding of active site features governing metal-oxo structure and reactivity.

KEYWORDS: heme enzyme, metal-oxo reactivity, hydrogen bonding, proton-coupled electron transfer, genetic code expansion, tryptophan analogue, cytochrome c peroxidase

Enzymes are the most proficient catalysts known, and consequently there is great interest in deciphering their sophisticated catalytic mechanisms. Site directed mutagenesis has been a staple technique in biochemistry for several decades as a means of elucidating the role(s) of key residues and molecular interactions.1 However, only a limited number of amino acid substitutions are possible as defined by nature’s genetic code. Under these constraints, substitutions designed to probe the importance of specific interactions (e.g., hydrogen bonds, π–π interactions) often lead to significant structural perturbations, making it difficult to parse out specific contributions to catalytic activity and complicating the interpretation of enzyme structure–activity relationships. This challenge is particularly acute when probing the role of the largest canonical amino acid, tryptophan, which has no close structural analogue within the genetic code. The availability of an expanded alphabet of amino acids provides a more surgical means of probing biological mechanisms by allowing substitutions of individual atoms or functional groups within proteins of interest.2–6 The power of this approach is exemplified through recent studies, whereby noncanonical cysteine and histidine analogues have been used to examine the role of axial heme ligands in controlling the reactivities of iconic ferryl intermediates compound I and compound II.7–13 These high-energy intermediates are the defining feature that drive catalysis across the entire family of heme enzymes, including P450s, peroxidases, nitric oxide synthases, and terminal oxidases.14,15 Consequently, there is great interest in understanding how different metal-oxo coordination environments within enzyme active sites control intermediate reactivity and overall catalytic function. Here, we use a noncanonical Trp analogue to examine directly the impact of hydrogen bonding interactions to the ferryl oxygen of compound I and compound II of cytochrome c peroxidase. Our data reveal how hydrogen bonding interactions are employed to control the reactivity of high-energy ferryl intermediates in enzyme active sites and thus advance our understanding of metal-oxo reactivity across a wide range of heme and nonheme iron enzymes.

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CeP employs a heme cofactor to reduce hydrogen peroxide in mitochondria using electrons from its biological redox partner ferrous cytochrome c (cyt). The reaction mechanism is comprised of three steps (Figure 1): (1) reaction of the resting ferric enzyme with hydrogen peroxide to generate compound I (CpdI), containing an oxidized ferryl (Fe(IV)=O) heme coupled to a neighboring Trp191 radical cation; (2) single electron reduction of CpdI by ferrous cyt to generate compound II (CpdII); and (3) single electron reduction of CpdII by a second equivalent of ferrous cyt. CpdII reduction is coupled with proton transfer to the ferryl oxygen, with the distal pocket His52 as the likely proton donor. In addition to redox active Trp191, CeP contains a second active site Trp51 whose N–H group forms a hydrogen bond to the ferryl oxygen of CpdI and CpdII (Figure S1). This interaction is also present in ascorbate peroxidase (APX), but is absent in many heme peroxidases, including the prototypical peroxidase from horse-radish, which contains a Phe residue in place of the Trp51 of CeP (Figure S1). Interestingly, Trp51Phe and Trp51Ala substitutions in CeP have been shown to substantially increase the rate of nonbiological oxidations of small molecule phenolics and anilines, along with more modest increases in cyt oxidation activity. The analogous Trp41Phe substitution in APX has been shown to increase activity with non-native phenolic substrates. However, the molecular origins of this increased reactivity of Trp51/41 variants of CeP and APX are not well understood. Some have argued that steric effects dominate, and that the introduction of smaller residues provides more space and flexibility in the distal heme pocket, which could give rise to the observed activity changes. Elsewhere, the increased reactivity of Trp51 variants of CeP has been ascribed to an increase in activation entropy, plausibly due to a more facile release of water from the heme iron. Others have suggested that hydrogen bonding between Trp51 and the ferryl oxygen has a stabilizing effect on key intermediates.

To resolve these uncertainties, we replaced Trp51 of Saccharomyces cerevisiae CeP with 3-benzothienyl-L-alanine (S-Trp), a close structural analog of tryptophan that cannot serve as a hydrogen bond donor, using an engineered pyrrolysyl-tRNA synthetase/pyrrolysyl-tRNA pair (PyrRS_S-Trp/tRNA\(^{\text{RS}}\)), which selectively incorporates S-Trp in response to the amber UAG stop codon. Stoichiometric replacement of the distal Trp51 residue with S-Trp was confirmed by MS analysis of the intact protein (Table S3). The X-ray crystal structure of CeP S-Trp (1.5 Å resolution, Table S2, Figure S2a) superimposes well with a previously reported CeP structure (Figure 2, PDB code: 2CYP, RMS deviation of 0.27 Å). Difference density (additional electron density) associated with the S atom of S-Trp was clearly visible. The geometry and environment of the heme cofactor and key active site residues are well preserved in the modified enzyme, with only minor conformational adjustments to the distal pocket His52, thus confirming Trp51S-Trp to be a highly conservative substitution. As anticipated, the X-ray crystal structure of the CeP S-Trp W191F double mutant is highly similar to that of CeP S-Trp (1.7 Å resolution, Table S2, Figure S2b), with a secondary structure superposed RMS deviation of 0.17 Å (Figure S2c).

Rapid mixing stopped-flow measurements were used to examine the nature of ferryl intermediates generated upon oxidation of CeP and CeP S-Trp with hydrogen peroxide. Consistent with previous studies, mixing CeP with 1.5 equiv of H\(_2\)O\(_2\) leads to spectral changes consistent with the formation of a neutral ferryl heme (Soret maxima at 420 nm and associated Q bands at 530 and 560 nm, Figure 3a). Similar spectral changes are observed upon oxidation of CeP S-Trp (Figure 3b), albeit with a slight decrease in the extinction coefficient of the Soret band (maxima at 420 nm) and Q-band features that are less well...
resolved than in the wild-type, suggesting that the CpdI state of CcP S-Trp is also comprised of a neutral ferryl heme. Time-dependent spectral changes were fitted to a sequential a → b model to derive rates for CpdI formation and are similar for both CcP and CcP S-Trp (k = 96 ± 3 and 100 ± 2 s⁻¹, respectively). To determine the identity of the protein radical in CcP S-Trp, CpdI was characterized by electron paramagnetic resonance (EPR). The CcP S-Trp CpdI EPR line shape is very similar to that of CcP CpdI (Figure 3c), confirming the formation of a coupled ferryl heme-Trp191 radical pair. The small change in the downfield “shoulder” g value from CcP to CcP S-Trp (2.041 to 2.033) likely arises due to the well-documented sensitivity of the magnetic coupling between the Trp191 cation radical and the ferryl heme to small local structural perturbations. To further characterize the ferryl intermediate upon disruption of the hydrogen bond to Trp51, resonance Raman spectra of CcP and CcP S-Trp were recorded in both the ferric and ferryl states (Figure 3d). Hydrogen bonding interactions to ferryl intermediates are thought to give rise to an increased Fe=O bond length and an associated reduction in Fe=O stretching frequencies. The intensity of the Raman feature associated with ferric CcP at 757 cm⁻¹ diminishes upon oxidation with H₂O₂, giving rise to a broad ferryl peak at 753 cm⁻¹, in accordance with the literature. Oxidation of CcP S-Trp also diminishes the 757 cm⁻¹ feature of the ferric state but instead leads to a new feature at 791 cm⁻¹ (Figure 3d), which we assign as the Fe=O stretch. The feature at 791 cm⁻¹ is also observed in the CcP S-Trp W191F double mutant (Figure S3). Density Functional Theory (DFT) models (vide infra) of the CcP ferryl state predict that the Trp51S-Trp substitution leads to a ~0.02 Å shortening of the ferryl bond, with an associated 44 cm⁻¹ increase in calculated Fe=O stretching frequency, which correlates well with the 38 cm⁻¹ increase observed experimentally (Figure S4).

To investigate the effect of the Trp51S-Trp substitution on ferryl heme stability, stopped flow measurements were repeated over a longer time frame. In wild-type CcP, the neutral ferryl heme was stable for >5 min (Figure S5c). In contrast, time-resolved UV/vis spectra reveal that the ferryl heme of CcP S-Trp decays to the ferric state with rate of k₃ = 0.04 s⁻¹ (Figure S5a). To gain insights into the origins of this reduced ferryl heme lifetime, we replaced the redox active Trp191 with Phe in CcP and CcP S-Trp, which allows the CpdI and CpdII states to be differentiated spectroscopically (Figure 4a). Previous studies have demonstrated that oxidation of CcP W191F generates a classical CpdI porphyrin π-cation radical state typical of most peroxidases with spectral features distinct to neutral ferryl heme systems. Oxidation of W191F variants of CcP and CcP S-Trp...
Trp with H$_2$O$_2$ (1.5 equiv) leads to the rapid (transient) formation of a CpdI porphyrin π-cation radical at similar rates in both variants ($k_1 = 126 \pm 1$ and $122 \pm 1$ s$^{-1}$, respectively, Figure 4b,c, Figures S6 and S7), as indicated by a substantial decrease in the Soret intensity (maxima at 406 and 407 nm, respectively). These data provide further evidence that Trp191 is the site of radical formation in both WT and CpP S-Trp. The CpdI states of CpCp W191F and CpCp S-Trp W191F subsequently decay to a neutral ferryl heme (CpdIII) with rates ($k_2$) of $24.2 \pm 0.1$ s$^{-1}$ and $6.93 \pm 0.01$ s$^{-1}$, respectively, indicating that the S-Trp substitution stabilizes the CpdI porphyrin π-cation radical (Figure 4b,c, Figures S6 and S7). In contrast, the S-Trp substitution decreases the lifetime of the CpdII state, which is stable for >5 min in CpCp W191F ($k_3 = \approx 0.003$ s$^{-1}$) but decays with a rate of $k_3 = \approx 0.07$ s$^{-1}$ in CpCp S-Trp W191F (Figure 4d, Figure S5b,d). This is similar to the observed increased reactivity of the neutral ferryl heme state of CpCp S-Trp vs wild-type CpCp.

To understand the contrasting impact of the S-Trp substitution on CpdI and CpdII reactivity in CpCp W191F, active site DFT calculations employing the Gaussian 09 software package were used to explore cluster models (see SI for details), which were generated based on a previously reported CpCp CpdII structure (PDB code: 5EJX). The calculations showed that replacement of Trp51 with S-Trp led to a modest 1.3 kcal mol$^{-1}$ reduction in the calculated electron affinity of CpdII, in accordance with the slower rate of CpdII reduction observed experimentally with the CpCp S-Trp W191F variant (Figure 4b). Our in silico results show that reduction of CpdII leads to spontaneous proton transfer from His52 (via an ordered water) to generate a ferric hydroxide state. Despite the increased reactivity of CpdII observed experimentally in CpCp S-Trp variants, PCET to CpdII is thermodynamically less favorable in CpCp S-Trp W191F ($\Delta G_{pcet} = 3.6$ kcal mol$^{-1}$, Figure 5, Table S4). We considered the possibility that the increased rate of CpdII decay could be attributed to single electron oxidation or sulfoxidation of S-Trp W191F. However, these off-pathway processes were discounted as (1) in silico sulfoxidation of S-Trp by CpdI, and to a greater extent CpdII, is endothermic (Table S12) and (2) prior studies have shown S-Trp to be considerably more difficult to oxidize to the radical state than Trp. To investigate the origins of the increased CpdII reactivity, we instead elected to deconvolute the PCET process into the component electron and proton transfer steps. We first calculated diabatic electron affinities for CpdII (EA$_{ii}$), which show that S-Trp substitution leads to a substantial reduction in electron affinity ($\Delta EA_{ii} = 10.0$ kcal mol$^{-1}$, Figure 5, Table S4). Similar $\Delta EA_{ii}$ values were calculated for adiabatic CpdII reduction by placing restrictions on the N–H/O–H bonds of Arg48, His52, and the ordered water. In contrast, proton transfer to the reduced CpdII states ($\Delta G_{h-transfer} = \Delta G_{pcet} – EA_{ii}$) is 6.4 kcal mol$^{-1}$ more favorable in CpCp S-Trp W191F. Numerous studies have demonstrated how the kinetics of PCET and related H atom transfers to ferryl centers are dominated by the basicity of the ferryl-oxygen and therefore we propose that the increased reactivity of CpdII observed experimentally upon S-Trp substitution can be ascribed to the substantial increase in $\Delta G_{h-transfer}$.

The combined experimental and computational data indicate that the Trp51 residue suppresses the reactivity and proton affinity of CpdII through hydrogen bonding to the ferryl oxygen. To understand the role of Trp51 in regulating the catalytic function of CpCp, we determined the kinetic parameters for the oxidation of ferrous cyt, and the nonbiological reductant guaiacol (ortho-methoxyphenol), by Trp51 and S-Trp51 variants of CpCp and CpCp W191F. The Trp51S-Trp substitution in CpCp and CpCp W191F causes dramatic 64-fold and 32-fold increases in the $k_{cat}$ of guaiacol oxidation, respectively, with only modest changes in $K_M$ (Figure 5, Figure S8a,b). This increased activity correlates well with the increased CpdII reactivity observed in the S-Trp containing variants. In contrast, the rate of cyt oxidation is only modestly affected by the Trp51S-Trp substitution ($k_{cat} = 819 \pm 46$ s$^{-1}$ and $596 \pm 33$ s$^{-1}$ for CpCp and CpCp S-Trp, respectively, Figure 5 and Figure S8c). As anticipated, replacement of the redox active Trp191 with Phe abolishes cyt oxidation activity in CpCp and CpCp S-Trp. Taken together, these data demonstrate how Trp51 tunes the lifetimes of key ferryl intermediates and works in synergy with the redox active Trp191 to control the substrate specificity of CpCp. Specifically, hydrogen-bonding from Trp51 extends the lifetimes of neutral ferryl heme intermediates, such as CpdII and the CpdII ferryl heme-Trp191 radical pair, which is likely important for biological function. This study builds upon previous work to improve the reactivity of CpCp toward small molecules by engineering binding sites for non-native substrates. What emerges is a complex picture whereby local ferryl coordination environments, the location and stability of key radical intermediates, and the presence of well-defined substrate binding sites work in synergy to control enzyme activity and selectivity. Nevertheless, the observation that Trp41Phe variants of APX (e.g., APEX2) also show increased activity with nonbiological substrates suggests that hydrogen-bonding to ferryl intermediates may control substrate specificity.
oxidations due to the formation of an inherently more reactive CpdII state. This study illustrates how an expanded genetic code can provide new tools to study complex bioinorganic reaction mechanisms. Genetically encoded cysteine and histidine analogues have been used by our lab and others to probe the influence of proximal heme ligands on CpdI and CpdII reactivity. Here, we have employed a noncanonical Trp analogue to elucidate how an active site hydrogen bond regulates Cp in modulating ferryl heme $p_K$, and reactivity. Proton-coupled electron transfers and related H atom transfers to high-energy metal-oxo intermediates are thought to be ubiquitous in biological systems. Consequently, we anticipate that the results presented will have wide-ranging implications for our understanding of metal-oxo reactivity in diverse enzyme active sites.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsau.1c00145. Materials and methods, calculations methods and data, MS data, UV–vis spectral data, kinetic analysis, structure statistics (PDF)

Accession Codes

The crystal structures of CpS-Trp and CcP S-Trp W191F were deposited in the RCSB Protein Data Bank (PDB) under accession numbers 6YT and 6Y2Y, respectively.

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The authors declare no competing financial interest.

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