Kinetic and Spectroscopic Studies of N694C Lipoxygenase: A Probe of the Substrate Activation Mechanism of a Nonheme Ferric Enzyme

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Abstract: Lipoxygenases (LOs) comprise a class of substrate activating mononuclear nonheme iron enzymes which catalyze the hydroperoxidation of unsaturated fatty acids. A commonly proposed mechanism for LO catalysis involves H-atom abstraction by an FeIII–OH site, best described as a proton coupled electron transfer (PCET) process, followed by direct reaction of O2 with the resulting substrate radical to yield product. An alternative mechanism that has also been discussed involves the abstraction of a proton from the substrate by the FeIII–OH leading to a σ-organoradical intermediate, where the subsequent σ bond insertion of dioxygen into the C–Fe bond completes the reaction. H-atom abstraction is favored by a high E° of the FeIII/FeII couple and a high pKₐ of water bound to the ferric state, while an organoradical mechanism would be favored by a low E° (to keep the site oxidized) and a high pKₐ of water bound to the ferric state (to deprotonate the substrate). A first coordination sphere mutant of soybean LO (N694C) has been prepared and characterized by near-infrared circular dichroism (CD) and variable-temperature, variable-field (VTVH) magnetic circular dichroism (MCD) spectroscopies (FeII site), as well as UV/vis absorption, UV/vis CD, and electron paramagnetic resonance (EPR) spectroscopies (FeIII site). These studies suggest that N694C has a lowered E° of the FeIII/FeII couple and a raised pKₐ of water bound to the ferric site relative to wild type soybean lipoxygenase-1 (WT sLO-1) which would favor the organoradical mechanism. However, the observation in N694C of a significant deuterium isotope effect, anaerobic reduction of iron by substrate, and a substantial decrease in kcat (~3000-fold) support H-atom abstraction as the relevant substrate-activation mechanism in sLO-1.

Introduction

The mononuclear nonheme iron enzymes comprise a large and expanding class of catalyzing reactions of medical, pharmaceutical and environmental significance. These enzymes utilize dioxygen as a cosubstrate and can be divided into two general groups: (1) dioxygen activating and (2) substrate activating enzymes. Oxygen activating mononuclear nonheme iron enzymes utilize a ferrous resting state which directly binds O2 to yield iron–oxygen intermediates which react with substrate to yield product. Substrate activating enzymes are characterized by their utilization of high-spin ferric sites to activate organic substrates for direct reaction with dioxygen. While numerous mononuclear nonheme enzymes exist, the majority are oxygen activating and differ only in the source of the additional reducing equivalents necessary for O2 activation (i.e., α-ketoglutarate-dependent, perlin-dependent, extradiol dioxygenases, Rieske dioxygenases). In contrast, lipoxygenases along with the intradiol dioxygenases are mononuclear nonheme ferric enzymes which activate substrate.

Lipoxygenases catalyze the regio- and stereospecific hydroperoxidation of 1,4-α,β-pentadiene-containing polyunsaturated carboxylic acids in plants and animals. In humans, lipoxygenases produce leukotrienes and lipoxins, signaling molecules that mediate conditions such as asthma, atherosclerosis, and psoriasis. Current insight into the molecular mechanism of lipoxygenase derives from a variety of structural, kinetic, and spectroscopic studies of soybean lipoxygenase (sLO-1) due to its ease of purification and stability. Crystallographic studies of ferrous sLO-1 have shown the active site iron to contain one C-terminal carboxylate ligand from Ile839, three histidine ligands (His690, His504, and His499), a water ligand, and a slow off rate of the oxygen binding site.

Along with other members of the family, soybean LO-1 (sLO-1) catalyzes the hydroperoxidation of fatty acids in a highly stereospecific manner. The stereochemical course of the reaction is determined by the nature of the substrate and the catalytic site. The reaction of 18:2 ω6 with soybean LO-1 has been studied in detail, and it is clear that the stereospecificity of the reaction is determined by the nature of the substrate and the catalytic site.

The reaction mechanism for lipoxygenase catalysis has been the subject of much debate. Early studies suggested that the initial step in the reaction is the abstraction of an H-atom from the substrate by the ferric enzyme, followed by direct reaction of O2 with the resulting substrate radical. This mechanism was supported by spectroscopic studies of the ferric enzyme, which showed a decrease in the E° of the FeIII/FeII couple and a raised pKₐ of water bound to the ferric state relative to wild type soybean lipoxygenase-1 (WT sLO-1) which would favor the organoradical mechanism. However, the observation in N694C of a significant deuterium isotope effect, anaerobic reduction of iron by substrate, and a substantial decrease in kcat (~3000-fold) support H-atom abstraction as the relevant substrate-activation mechanism in sLO-1.

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dense side chain carbonyl of Asn694, at 2.87 Å. 9,10 Circular dichroism (CD) and magnetic circular dichroism (MCD) studies of resting sLO-1 provided further insight into the active site structure having shown it to be a 4/60% mixture of five-coordinate (5C) and six-coordinate (6C) ferrous species which becomes 6C upon substrate binding.11 The difference in coordination is due to the coordination flexibility of the Asn694 ligand which contributes to the reactivity of the site.

For activity, the enzyme must be oxidized to the ferric state. A commonly proposed mechanism for sLO-1 is shown in Scheme 1A. The catalytic cycle begins with hydrogen atom abstraction of the pro-S hydrogen from the 1,4-cis,cis-pentadiene fatty acid substrate by the FeIII-OH species,12 shown to be a 5C species by electron paramagnetic resonance (EPR) with the signal being axial in a nonheme environment,13 to form a pentadienyl radical and the FeII-OH intermediate.6,13,15,16 A large kinetic isotope effect (KIE) has been reported for sLO-1 (KIE 80)17 and human 15-LO-1 (KIE 50)18,19 indicating that C-H cleavage contributes to the rate-determining step and its mechanism is via hydrogen tunneling. For this mechanism, the coordination flexibility (6C FeII 5C FeIII) attributed to the Asn694 ligand, has been shown by site-directed mutagenesis to play an important role in tuning the active site for catalysis.20 Density functional theory calculations on H-atom abstraction by the FeIII-OH site of sLO have suggested that the coordination flexibility contributes to the reaction energetics by stabilizing the FeII relative to the FeIII state raising the redox potential and the pKs of the water ligand of the reduced site (to favor H-atom abstraction).21,22 The transition state for H-atom abstraction is best described as a proton coupled electron transfer (PCET) process as the electron tunnels directly from the substrate to the iron through a superexchange pathway involving an FeO-H-bridge. Dioxygen would react directly with the substrate radical formed in the PCET process to yield a peroxyl radical which subsequently oxidizes the FeIII-OH to give the active FeIII-OH- substrate and the hydroperoxide product.

Since both the E° of the FeII/FeIII couple and pKs of water bound to the ferrous site are affected by the strength of the ligands to FeII, first coordination sphere mutants have been studied to gain further insight into the mechanism of catalysis in LOs. These mutant studies have focused on the Asn694 ligand responsible for the coordination flexibility of sLO-1 shown to be important in tuning the site in sLO-1 for catalysis. Two such mutants, N694H and N694G, were previously generated and characterized.23,24 In contrast to wild type soybean lipoygenase-1 (WT sLO-1) which is 6C in the ferrous state with the substrate analogue bound and converts to a 5C site upon oxidation, CD and MCD studies showed that both N694H and N694G were 6C in both the reduced and oxidized forms. Furthermore, detailed analysis of both mutants indicated that they had a lowered E° relative to WT sLO-1 due to the stronger donor ability of the bound ligand compared to Asn (His for N694H and hydroxide for N694G). The lowered E° correlated with a decrease in kcat for the mutants, indicating that the E° of the FeII/FeIII couple is a key driving force for H-atom abstraction.

To gain further molecular-level insight into the factors controlling catalysis in LO, we have extended our first sphere mutant studies of sLO-1 to a cysteine mutant of Asn694 (N694C). The thiolate ligand of cysteine in the mutant would have a higher affinity for the oxidized site which would lower the potential of the FeII/FeIII couple, and the Cys is a strong donor ligand which would decrease the inductive effect of the FeIII on the coordinated water, raising the its pKs. Interestingly, this perturbation of the iron site would, in principle, favor an alternative, organoiron mechanism (Scheme 1B). In this mechanism, initial abstraction of a proton by the enzyme is facilitated by complexation of the carbanion to FeIII, yielding a σ-organooiron intermediate. The subsequent bond insertion of dioxygen in the C=FeIII bond would complete the reaction to yield product. Importantly, several aspects of this organoiron mechanism have been suggested to be feasible by model studies, where reaction of allylic organoiron intermediates with dioxygen was shown to be a facile process, able to compete with homolysis cleavage of the C=FeIII bond.26 In contrast to H-atom abstraction, the σ-organooiron mechanism would be favored by a low E° to keep the site oxidized and high pH for water bound to the ferric state to deprotoenate the substrate. This organooiron mechanism has not been directly experimentally evaluated.

Near-infrared CD and variable-temperature variable-field (VTTH) MCD studies are utilized to determine the geometric and electronic structure of the ferrous form of N694C LO, and...
Materials and Methods

Linoleic acid (LA) was purchased from Aldrich Chemical Co. and perdeuterated linoleic acid (D-LA) was purified from a mixture of perdeuterated algal fatty acids from Cambridge Isotope Labs, as previously described. All commercial fatty acids were repurified using a Higgins Semi-Prep Haisil (25 cm × 4.6 mm) C-18 column. Solution A was 99.9% MeOH and 0.1% acetic acid, while solution B was 99.9% H2O and 0.1% acetic acid. An isotropic elution of 85% A/15% B was used to purify all fatty acids, and they were stored at −80 °C for a maximum of 6 months. All other chemicals were reagent grade or better and were used without further purification.

Mutagenesis, Overexpression, and Purification of N694C. Site-directed mutagenesis, overexpression, and purification of WT sLO-1 and N694C followed a protocol outlined previously. In brief, following expression of the protein in BL21-DE3 (Escherichia coli), the cells were harvested by centrifugation and their membranes were disrupted by sonication. Cell debris was pelleted, and the supernatant was first dialyzed against 20 mM Bis-Tris buffer (pH 6.0) and then applied to an SP-Sephadex high-flow ion exchange column (Pharmacia), which was equilibrated with the same buffer. Eluted fractions containing lipoxygenase activity were pooled, concentrated, dialyzed against 20 mM Bis-Tris buffer (pH 6.0), and applied to a Macro-Prep 25-S ion exchange column (Bio-Rad). After concentration and buffer exchange into 0.1 M Borate (pH 9.2), the isolated enzyme was estimated to be greater than 90% pure (SDS-PAGE). Iron content of all lipoxygenase enzymes were determined on a Finnegan inductively coupled plasma mass spectrometer (ICP-MS) using internal standards of Co²⁺-EDTA, and data were compared with those of standardized iron solutions.

Steady-State Kinetic Measurements. Steady-state kinetic values were determined by following the formation of the conjugated product at 234 nm (ε = 2.5 × 10⁴ M⁻¹ cm⁻¹) with a Perkin-Elmer Lambda 40 spectrophotometer. No photodegradation of the product was observed. All kinetic measurements were standardized to iron content. The assay samples were 2 mL in volume with substrate concentrations ranging from 1 to 80 μM (Borate buffer, pH 9.2, 25 °C) under constant stirring with a rotating magnetic bar. The total ionic strength was adjusted to 50 mM by addition of NaCl. The temperature was controlled using a circulating water bath and jacketed cuvette system. Stock solution of LA was prepared in 95% ethanol and diluted into buffer so that the total ethanol concentration was less than 1.5%. Fatty acid concentrations were verified by allowing the enzyme reaction to proceed to completion and quantitating the conjugated product at 234 nm. Initial rates (up to the first 15% of the reaction) for each substrate were fitted to the Michaelis–Menten equation using the KaleidaGraph (Synergy) program and errors determined.

Competitive Kinetic Isotope Effect. The kinetic isotope effect (KIE) reactions were performed as previously described. Briefly, a protoi/ perdeutero LA mixture, of known molar ratio, was reacted with LO in the kinetic buffer described above. The reaction was monitored at 234 nm and stopped with an acid quencher at less than 5% total LA consumption. The acidified reaction mixture was extracted with methylene chloride, evaporated to dryness, reconstituted in methanol, and injected onto an RP-HPLC column (Higgins Analytical, 250 mm × 4.6 mm). The products eluted at 1 mL/min (isocratic mobil phase 74.9% methanol, 25% water, 0.1% acetic acid). The molar protoi/perdeutero 13-(S)-hydroperoxy-9,11-(Z,E)-octadecadienoic acid (the oxidation product of LA) (HPOD) ratios were equated to the corresponding peak area ratios and the KIE calculated as described previously.

UV–vis Absorption and CD spectroscopies were carried out at 4 °C. The UV–vis absorption spectra were recorded on a Perkin-Elmer Lambda 40 monochromometer. Spectra were taken in 0.1 M Borate (pH 9.2) at 0.1 M ionic strength (adjusted with NaCl) at 4 °C. CD spectra were obtained using an Aviv 60DS spectropolarimeter in 0.1 M Borate (pH 9.2). CD spectra were baseline corrected by subtracting the buffer and cell background from the raw data. Titrations were performed by the addition of 1 equiv of HPOD to 0.6 mL of lipooxygenase (20 mg/mL).

EPR Spectroscopy. EPR measurements were performed on a Bruker ESP-500E spectrometer equipped with a dual mode X-band cavity and an Oxford Instruments ESR-900 helium flow cryostat. The program HAM (Dr. M. Hendrich, Carnegie Mellon University, Pittsburgh, PA) was used for data manipulation and EPR spectral simulation, as described previously. In brief, the signals were fit with an effective S' = 1/2 model which uses a g-dependent line width that properly accounts for field-dependent transition probabilities. The temperature dependence was also fit with this program using the standard Boltzmann equation. The amount of EPR-active FeI in the protein samples and the temperature dependence were determined relative to an FeI-EDTA standard. A 100 μM Fe solution was made from a 1000 ppm iron standard (Fischer Scientific), with addition of a 10-fold excess of EDTA in 25% glycerol. The EPR spectra were recorded at a modulation frequency of 100 kHz and at microwave frequencies of about 9.44 GHz. Precise microwave frequencies were recorded for individual spectra to ensure accurate g-alignment. Other EPR parameters are specified in the figure legends.

Anaerobic Reactivity and Spectroscopic Properties of Ferric N694C. A ferrous sample of N694C (108 mg/mL in the kinetic buffer) was titrated anaerobically with 1 equiv aliquots of HPOD until the ferric form of N694C was fully generated (as seen by the 762 nm band intensity, vida infra). An EPR sample was removed and determined to manifest a rhombic ferric signal. LA substrate (1 equiv) was then added anaerobically, and the decrease in signal intensity at 762 nm was measured. An EPR sample of the substrate oxidized species with LA added was removed, and its change was monitored with EPR.

Near-IR CD and MCD Spectroscopy. Near-infrared (600–2000 nm) CD and MCD data were recorded on a Josco J-200D spectropolarimeter with a liquid N₂-cooled InSb detector and equipped with an Oxford Instruments SM4000-7T (T) superconducting magnet/cryostat capable of fields up to 7 T and temperatures down to 1.5 K. The N694C CD sample (~1.5 mM) was exchanged into deuterated CHES buffer (50 mM, pH 9.2) and kept at 5 °C during data collection with a circulating cooling bath attached to the sample holder. Glycerol-d₁ was added at 60% (v/v) to the N694C/FeII solution for preparation of the MCD sample, and CD spectra were taken with and without glycerol addition to ensure that the site was unaffected by the glassing agent (Note: no change in the CD spectrum due to glycerol was observed). CD spectra were corrected for buffer and cell baselines by subtraction. Low-temperature MCD data were obtained in custom-made cells consisting of two Infrasil quartz disks separated by a 0.3 cm thick neoprene O-ring spacer into which the sample was injected and secured between two copper plates. The MCD spectrum was corrected for the natural CD and zero-field baseline effects caused by strain in the glasses by averaging the positive and negative field data at a given temperature (i.e., (T T − (−)T/2)). VTVH data were normalized to the maximum observed intensity over all isotherms for a given wavelength, and the
Results

Protein Purification and Kinetic Determinations. N694C and WT sLO-1 were purified with yields of 2–4 mg/L, with metal contents of 55 ± 10% and 80 ± 5%, respectively.

The steady-state parameters of N694C sLO, $k_{\text{cat}}$ (0.098 ± 0.002 s$^{-1}$), $k_{\text{cat}}/K_M$ (0.026 ± 0.002 s$^{-1}$ µM$^{-1}$), and $K_M$ (3.8 ± 0.3 µM) (Table 1), were determined at 22 °C ($k_{\text{cat}}$, the rate constant for product release; $k_{\text{cat}}/K_M$, the rate constant for substrate capture; $D_{k_{\text{cat}}}/K_M$, the competitive primary kinetic isotope effect). The $k_{\text{cat}}$ value of N694C was ∼2900 times lower than that of the WT sLO-1, and the $k_{\text{cat}}/K_M$ value was ∼730 times lower; however, the $K_M$ value is similar to that observed for WT sLO-1. The competitive kinetic isotope effect data ($D_{k_{\text{cat}}}/K_M$) for N694C were temperature-independent, with values of 44 ± 2 at 15 °C, 48 ± 3 at 22 °C, and 47 ± 6 at 30 °C (average value of approximately 46 ± 2). In previous studies, it was shown that WT sLO-1 displayed a temperature-dependent, competitive $D_{k_{\text{cat}}}/K_M$ at low substrate concentration (5 µM). This temperature dependence is due to the multiple rate-limiting steps in the mechanism: substrate diffusion, hydrogen bond rearrangement, and hydrogen bond abstraction (C–H bond cleavage). However, for N694C, the $D_{k_{\text{cat}}}/K_M$ is temperature-independent, indicating that the rate-determining step for N694C is solely limited by the C–H bond cleavage since tunneling mechanisms are not temperature-dependent. The data for N694C are comparable to our previous data for N694H, which manifested a $D_{k_{\text{cat}}}/K_M$ of 60 ± 9 at 30 °C. Although the magnitude of $D_{k_{\text{cat}}}/K_M$ for N694C is less than that observed for N694H, the magnitude of the $D_{k_{\text{cat}}}/K_M$ is much larger than semiclassical theory predicts (KIE ~7) and is consistent with a tunneling mechanism for hydrogen atom abstraction.

Spectroscopy. (1) Fe$^{III}$ Site. Oxidation of N694C was accompanied by an increase in absorbance at 342 nm (ε = 2650 M$^{-1}$ cm$^{-1}$) and 762 nm (ε = 760 M$^{-1}$ cm$^{-1}$) (Figure 1, samples corrected for iron content). The latter transition is characteristic of a Cys to Fe$^{III}$ ligand-to-metal charge-transfer band. It is lower in energy and has less intensity than the Cys to Fe$^{III}$ charge transfer observed, for example, in superoxide reductase.33,34 UV/vis CD spectroscopy manifests a band at 345 nm (Δε = 2.2 M$^{-1}$ cm$^{-1}$) and a shoulder at 420 nm (Figure 2). The Δε value was corrected for iron content and is comparable in magnitude to that of the WT sLO-1 (Δε = 1.45 M$^{-1}$ cm$^{-1}$), assigned as a His to Fe$^{III}$ charge-transfer transition.

The EPR spectrum of as-isolated N694C (60 µM iron) in frozen solution at 4 K exhibited a weak signal at g ~4.3 (data not shown). The low intensity of this signal represents less than 5% of the total iron and indicates that the enzyme is mostly in its ferrous state. Addition of 1 equiv of HPOD (the hydroperoxide product of sLO) to ferrous N694C oxidized the iron to its ferrous state. Addition of 1 equiv of HPOD (the hydroperoxide product of sLO) to ferrous N694C oxidized the iron to the catalytically competent ferric species. The ferric N694C at pH 9.2 (0.1 M borate, 0.1 M ionic strength) exhibited a complex multicomponent EPR signal, centered around g ~4.3, with a minor species at g ~6 (Figure 3A). The ferric species was also generated at pH 7.2 (0.1 M Tris, 0.1 M ionic strength), and only slight perturbations were observed.

The quantification of the amount of Fe$^{III}$ contributing to the multicomponent signal was complicated by the fact that the signal is spread over a large field range (~2300 G). This problem was addressed by the EPR spectral simulations (see Supporting Information). The simulated spectra were integrated over the entire field range to obtain the total EPR intensity from transitions within the middle Kramers doublet for the rhombic signal and the $M_S = ±1/2$ for the axial signal (see Figure 3B and Supporting Information). Utilizing the simulated ($g_{av}$ corrected) intensity, it was determined that greater than 85% of the expected ferric signal was observed and that the relative

Table 1. Comparison of Kinetic Parameters for WT and N694C sLO-1

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<th>$k_{\text{cat}}$ (s$^{-1}$)</th>
<th>$k_{\text{cat}}/K_M$ (µM$^{-1}$ s$^{-1}$)</th>
<th>$K_M$ (µM)</th>
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<tr>
<td>WT</td>
<td>287(5)</td>
<td>19(1)</td>
<td>15(1)</td>
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<tr>
<td>N694C</td>
<td>0.098(0.002)</td>
<td>0.026(0.002)</td>
<td>3.8(0.3)</td>
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* Numbers in parentheses represent errors as determined from replicate determinations. * Numbers were collected at 22 °C, in parallel.
concentrations of the EPR species for ferric N694C (pH 9.2) were 80 ± 8% for the rhombic signal and 20 ± 2% for the axial signal.

An as-isolated, reduced N694C sample was oxidized with 1 equiv of HPOD, anaerobically. An increase at 762 nm and an increase of the rhombic EPR signal were observed. Anaerobic LA substrate (1 equiv) was subsequently added, and the rate of the disappearance of the 762 nm band was observed to be 0.017 ± 0.003 s⁻¹ (Figure S2). This value is consistent with the rate of enzymatic turnover, considering that substrate is not saturating. The EPR signal of the ferric N694C disappeared upon substrate addition (paralleling the Cys→FeIII charge-transfer band in absorption), indicating reduction of the ferric species to be associated with the anaerobic reaction of the enzyme with substrate.

(2) FeII Site. Near-infrared CD and MCD spectroscopies were employed to study the high-spin ferrous active site in N694C in solution as analysis of the energies and splitting pattern of CD/MCD bands provides information about the geometric and electronic structure of the ferrous active site. The 278 K CD spectrum of N694C/FeII contains a single negative ligand field band at 6150 cm⁻¹ (Figure 4A), CD spectra measured both without and with the glassing agent glycerol present were unchanged. The 1.6 K, 7 T MCD spectrum of N694C/FeII (Figure 4B) contains a large positive ligand field feature at ~6050 cm⁻¹ and a weak negative feature at ~11 450 cm⁻¹. The saturation magnetization behavior for N694C/FeII collected at 6250 cm⁻¹ (Figure 4B, inset) is well-fit by a positive ZFS non-Kramers doublet model (i.e., large nesting) with ground-state spin Hamiltonian parameters of $D = 12.5 ± 0.3$ cm⁻¹ and $|E| = 2.6 ± 0.2$ cm⁻¹, corresponding to $\Delta = +500 ± 150$ cm⁻¹ and $|V/2\Delta| = 0.29 ± 0.02$ which reflect the axial ($E_{ax} - E_{xy} = \Delta$) and rhombic ($E_{ax} - E_{xy} = V$) dσ splitting of the $t_{2g}$ set of d orbitals on the FeII. The observed CD and MCD excited-state splittings and energies of the two transitions of N694C/FeII are consistent with a square pyramidal 5C ferrous site. It should be noted that Fe K-near-edge X-ray absorption analysis performed on N694C/FeII indicates that the cysteine sulfur is not bound to the FeII site and, thus, is the ligand lost to generate the 5C site.36

Discussion

A first sphere mutant of sLO-1 (N694C) has been prepared in which the Asn694 ligand responsible for the coordination flexibility observed in WT sLO-1 has been selectively mutated to a Cys. A combination of CD and VTVH MCD spectroscopies indicates that the FeII site of N694C is a 5C square pyramidal species in contrast to the 40/60% mixture of 5C and 6C ferrous species in WT sLO-1/FeII. The Cys residue is not coordinated to the FeII site of the mutant, but oxidation of the site leads to ligation of the cysteine sulfur to the FeIII based upon the presence of a Cys→FeIII charge-transfer band at 762 nm ($\epsilon = 760$ M⁻¹ cm⁻¹).

The observed Cys→FeIII charge-transfer band in N694C is at lower energy with decreased intensity compared to the Cys→FeIII transition in WT sLO-1/FeIII in such as superoxide reductase.33,34 One possible contribution to the decreased intensity of the transition could be partial coordination of the ferric site by Cys. However, from EPR of ferric N694C, the rhombic signal, indicative of a 6C site in sLO where the Cys S occupies the sixth coordination site, is present at 80%. Correcting for this still leads to a factor of ~2.5 lower extinction coefficient relative to superoxide reductase ($\epsilon$ (SOR at 660 nm) = 2500 M⁻¹ cm⁻¹).34 However, a weakened FeIII→S bond in N694C compared to superoxide reductase would both decrease the intensity of the transition and shift it to lower energy as observed. A weaker Cys→FeIII bond could reflect the fact that the cysteine side chain is one carbon shorter than the Asn ligand.


it replaced. This is also consistent with the loss of the Cys ligand upon reduction of the site.

Previous studies of WT sLO-1 have shown that it is converted from a 6C FeII site upon substrate analogue binding to a 5C FeIII site.12,14 In contrast, the present study indicates that N694C sLO-1 is converted from a 5C FeII site with a substrate analogue bound (oleic acid, see Supporting Information) to a 6C FeIII site as determined from EPR data which show a predominately rhombic site indicative of 6C.37 The FeIII form of N694C is the catalytically active species analogous to WT sLO-1 as determined by UV/vis absorption and EPR studies of the decay of this species at a catalytically relevant rate in the anaerobic reaction with substrate. Importantly, the $k_{cat}$ for N694C is significantly reduced (~3000-fold) relative to WT sLO-1, and the iron active site in the mutant is anaerobically reduced upon substrate addition.

Our spectroscopic studies of N694C support the presence of an iron active site for N694C with a lowered $E^o$ of the FeII/FeIII couple compared to WT sLO-1 due to the presence of a 6C ferric site, with the cysteine sulfur coordinated in the ferric but not ferrous state. This stabilizes the FeIII site relative to the FeII site. The $pK_a$ of the coordinated water in the 6C ferric site of the mutant should be increased, as the site is 6C and the cysteine donor ligand would tend to decrease the donation of coordinated $H_2O$ to FeIII. Thus, the Asn694 $\rightarrow$ Cys694 mutation results in an iron active site with an $E^o$ of the FeII/FeIII couple and a $pK_a$ of water bound to the ferric site that would, in principle, be more effective than WT sLO-1 in a $\sigma$-organoiron mechanism. Alternatively, with respect to H-atom abstraction, the coordination change relative to WT sLO-1 (loss of the cysteine ligand to produce a 5C FeII site) would both lower $E^o$ of the FeII/FeIII couple and lower the $pK_a$ of water bound to the ferrous site which would disfavor this reaction. In fact, the enzymatic turnover data are consistent with an H-atom abstraction mechanism for N694C due to the observation of a significant deuteration isotope effect on rate, a substantially reduced $k_{cat}$ (~3000-fold) of N694C relative to WT sLO-1, and the observed anaerobic reduction of iron by substrate. The iron active site would remain in the ferric state throughout the catalytic cycle in a $\sigma$-organoiron mechanism.

In addition to $E^o$ and $pK_a$, possible structural changes due to the Asn694 $\rightarrow$ Cys694 mutation could contribute to the decreased reactivity. Since the cysteine sulfur is not coordinated in N694C/FeIII, the ligands likely remain unchanged relative to the WT sLO-1/FeII 5C component (3His, 1 C-terminal carboxylate from Ile839 and $H_2O$). However, a perturbation of the 5C ferrous site of N694C is observed compared to the 5C site in WT sLO-1/FeII (Figure 4B,C) that both excited-state transitions have shifted to higher energy and the sign of the higher energy transition has become negative.12 In addition, the splitting of the $T_{2g}$ ground-state obtained from the VTVH MCD data in Figure 4B (inset) has decreased in the mutant. Previously it has been shown that the N694 ligand is involved in a H-bonding network with residues Q697 and Q495 in the second sphere, linking the substrate binding pocket to the flexible N694 ligand and, thus, playing a role in the coordination flexibility of this residue (Figure 5).11 Binding of substrate, glycerol, or inhibitor (oleic acid) affects the H-bonding, resulting in conversion to a purely 6C site.20 However, the N694C mutation eliminates the possibility of a H-bond from residue 694 to Q697, supported by CD studies showing no perturbations of the FeII due to glycerol or substrate analogue oleic acid binding (see Figure S3). Crystallographic studies of other first sphere mutants (N694H and N694G)11,24 which also lack H-bonding from residue 694 to Q697 indicate that perturbing this network also affects the interactions of these second sphere residues with the coordinating H499 ligand and, thus, its interaction with the FeII. Therefore, the changes in the MCD spectrum of N694C/FeII relative to WT sLO-1/FeII are consistent with the perturbation of the H-bonding network and the effects of this on the H499 ligand. However, since both N694C/FeII and WT sLO-1/FeII are 5C with the same ligation, this structural perturbation should not have a significant effect on $E^o$. For the catalytically relevant ferric site of N694C sLO-1, the structural difference from WT sLO-1/FeII is the ligation of Cys which could sterically affect substrate binding. However, the fact that the $K_M$ value for the N694C mutant is similar to that for WT sLO-1 and to that previously reported for another first sphere mutant, N694H ($K_M$ ~4 $\mu$M),23 indicates that the mutation results in only minor changes in the substrate binding pocket.

Finally, based on crystallographic studies of WT sLO-1 in the absence of substrate, it was proposed that the presence of certain active site residues such as Gln495 and Gln697 restrict access to the Fe site to disfavor formation of the $\sigma$-organoiron intermediate.10 However, the possibility of structural rearrangement of the protein active site to accommodate this species could not be eliminated. Subsequent crystallographic studies do indicate the possibility of structural changes in some of these residues upon substrate binding.11 However, the spectroscopic and kinetic data presented here argue against a $\sigma$-organoiron intermediate.

In summary, a first sphere mutant of sLO-1, N694C, has been prepared and characterized which allows evaluation of the $\sigma$-organoiron mechanism previously proposed for the reaction of lipoxygenases. While the mutant has a lowered $E^o$ of the FeII/FeIII couple and a raised $pK_a$ of water bound to the ferric site which would favor a $\sigma$-organoiron reaction, the kinetic data of N694C show a significant deuteration isotope effect on rate, an observed anaerobic reduction of iron by substrate, and a substantially reduced $k_{cat}$ (~3000-fold) of N694C relative to WT sLO-1. These results experimentally evaluate the organoiron mechanism and provide further support for H-atom abstraction as the catalytically relevant mechanism in lipoxygenase.
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Supporting Information Available: EPR simulation and integration for ferric N694C, LA oxidation by N694C/FeIII monitored by absorption and the 278K CD spectra of N694C/FeII+ 60% (v/v) glycerol and N694C/FeII+ oleic acid (saturating). This material is available free of charge via the Internet at http://pubs.acs.org.