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# Geometric Structure Determination of N694C Lipoxygenase: A Comparative XAS Near-Edge and EXAFS Study

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### Abstract

The mononuclear non-heme iron active site of N694C soybean lipoxygenase (sLO1) has been investigated in the resting ferrous form using a combination of Fe K-pre-edge, near-edge (using MXAN full multiple-scattering approach) and EXAFS methods. The results indicate that the active site is six coordinate (6C) with a large perturbation in the first shell bond distances in comparison to the more ordered octahedral site in wild type sLO1. Upon mutation of the asparigine to cystiene, the short Fe-O interaction with asparigine is replaced by a weak Fe-(H<sub>2</sub>O) which leads to a distorted 6C site with an effective 5C ligand field. In addition, it is shown that near-edge multiple scattering analysis can give important three-dimensional structural information, which usually cannot be accessed using EXAFS analysis. It is further shown that relative to EXAFS, near-edge analysis is more sensitive to partial coordination numbers and can be potentially used as a tool for structure determination in a mixture of chemical species.

## 1. Introduction

Extended X-ray absorption fine structure is a powerful technique for local structure determination on systems in both solution and solid (amorphous and crystalline) states.<sup>1–3</sup> Traditionally EXAFS has been applied to a broad range of scientific fields, especially for systems that cannot be obtained in the crystalline form for X-ray diffraction measurements. The EXAFS region typically extends from ~50 to 1500 eV above the edge inflection of the Xray absorption spectrum, however, the magnitude of the EXAFS signal is reduced rapidly with energy. This imposes potential limitations, for example, i) at low absorber concentration, sufficient signal-to-noise at higher energies may not be obtained, constraining the data range, *ii*) photoreduction and beam damage may impair multiple scan averaging for low-concentration samples or, *iii*) inherently weak EXAFS signals, which inhibit data collection to high energies. These limitations are in particular applicable to non-heme iron proteins, which usually have irregular first shell (N/O) ligation and cannot be obtained at higher concentrations.<sup>3</sup> The nearedge region of an XAS spectrum (~-10 to 200 eV) successfully overcomes the concentration and weak signal limitations in such systems. Also, since good quality near-edge data can be collected significantly faster, beam damage and photoreduction can be minimized. The nearedge fitting methodology implemented into the MXAN code (Minuit X-ray Absorption Near-

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Edge) which has been developed by Benfatto *et al.*, has been successfully applied to several small inorganic molecules.<sup>4–7</sup> In a recent study, the full multiple scattering theory implemented in MXAN was applied to  $[Cu(TMPA)(OH_2)](ClO_4)$ , which is a model complex that mimics the active site of cytochrome *c* oxidase (*CcO*).<sup>8</sup> It was shown that MXAN was successful in simulating the near-edge spectrum and in describing the complex multiple-scattering from the TMPA ligand system. In addition, it was shown that the multiple scattering approach is sensitive to small structural and angular variations.

Lipoxygenases (LO's),<sup>9–13</sup> which are non-heme iron containing enzymes with mixed N/O donor ligands, are systems challenged by the abovementioned EXAFS limitations. Although two different crystal structures of soybean lipoxygenase (sLO) have described the resting ferrous active site as six-coordinate (6C) (three histidine ligands (His), the C-terminal isoleucine (Ile) ligated to the Fe with the free carboxylate oxygen, H<sub>2</sub>O and a weakly coupled asparigine (Asn) ligand)<sup>14,15</sup> and/or strongly distorted four-coordinate (4C) (considering only the four strongly coordinating first shell ligands, i.e. three His and the free carboxylate oxygen of the C-terminal Ile),<sup>16</sup> a combination of CD (circular dichroism) and MCD (magnetic circular dichroism) studies,<sup>17</sup> which is a powerful technique to identify the coordinate (5C) and six-coordinate (6C), in a 40/60 ratio which becomes 6C upon glycerol addition. This has been confirmed by EXAFS measurements on WT sLO in the presence of 30% glycerol.<sup>17</sup>

In this study, the MXAN near-edge multiple-scattering method has been used, at the Fe K edge, in combination with pre-edge and EXAFS analysis to study the ferrous form of the N694C mutant of lipoxygenase. In this mutant, the weakly coordinating first-coordination sphere amino acid Asn694 has been replaced by a potentially strongly coordinating cysteine ligand. CD and MCD studies indicate that the ferrous site is 5C and remains 5C upon glycerol binding. <sup>18</sup> However, in the absence of a crystal structure there are several structural possibilities for a 5C site. These structural possibilities (elaborated in the Results and Analysis section) have been evaluated using this combined XAS analysis approach. The results indicate that the active site is distorted 5+1C with a weakly coordinated axial H<sub>2</sub>O at an Fe-O distance of 2.5 Å, and that this weak coordination of the water molecule to the ferrous center retains an approximately 5C ligand field (in the presence of glycerol). It is shown that the near-edge multiple scattering approach can differentiate between structural models proposed by EXAFS and can be used as a powerful structural tool if adequate EXAFS data are not available.

#### 2. Experimental Section

#### 2.1 Sample Preparation

Site-directed mutagenesis, overexpression, and purification of N694C sLO followed a protocol outlined previously.<sup>19</sup> N694C was purified with a yield of 2–4 mg/L, with a metal content of  $55 \pm 10\%$ . N694C was concentrated and 30% (v/v) glycerol was added to form an optical quality glass with a final Fe<sup>II</sup> concentration of ~ 1 mM. The enzyme sample was then transferred under an inert atmosphere wet-box to an XAS sample cell and frozen immediately in liquid nitrogen.

#### 2.2 X-ray Absorption Spectroscopy

The X-ray absorption spectra of N694C mutant of LO were measured at the Stanford Synchrotron Radiation Laboratory on the focused 16-pole 2.0 T wiggler beam line 9-3 and the unfocussed 8-pole 1.8 T wiggler beam line 7-3 under standard ring conditions of 3 GeV and 60–100 mA. A Si(220) double crystal monochromator was used for energy selection. A Rh-coated harmonic rejection mirror and a cylindrical Rh-coated bent focusing mirror were used for beam line 9-3 whereas the monochromator was detuned 50% at 7998 eV on beam line 7-3

Ge solid-state detector windowed on the Fe K $\alpha$  signal. Internal energy calibration was accomplished by simultaneous measurement of the absorption of an Fe-foil placed between two ionization chambers situated after the sample. The first inflection point of the foil spectrum was assigned to 7111.2 eV.

Data represented here are a 21-scan average spectrum, which was processed by fitting a secondorder polynomial to the pre-edge region and subtracting this from the entire spectrum as background. A three-region spline of orders 2, 3 and 3 was used to model the smoothly decaying post-edge region. The data were normalized by subtracting the cubic spline and by assigning the edge jump to 1.0 at 7120 eV using the *SPLINE* program in the XFIT suite of programs.<sup>20</sup>

#### 2.3 Pre-Edge Data Analysis

Least squares fit were performed using the  $EDG_FIT$  program<sup>21</sup> to quantify the intensity and energy of the pre-edge feature. The pre-edge features were modeled using 50:50 Lorentzian:Gaussian pseudo-Voigt functions. The rising-edge background was modeled using a pseudo-Voigt function, which mimicked the white line associated with the edge transition. Additional pseudo-Voigt peaks were also used to mimic shoulders on the rising edge. The data were fit over several different energy regions. The least-square error and a comparison of the second derivatives of the data and fit were used to determine the goodness of fit. The standard deviations in energy and intensity over all successful fits were used to quantify errors in these parameters.

#### 2.4 Near-Edge Data Analysis

The near-edge simulations were performed using MXAN,<sup>4,7</sup> which performs a full multiplescattering analysis of the data to obtain structural information.<sup>7,22</sup> The following programs are integrated in MXAN: VGEN, a generator of Muffin-tin potential, the CONTINUUM code for full multiple-scattering cross-section calculation and the MINUIT routines of the CERN library for parameter optimization. The starting structures used in the simulations were generated by modifying the crystal structure of wild-type sLO.<sup>15</sup> Several starting structures were considered which included both five-coordinate (5C) and six-coordinate (6C) Fe centers. The data were fit over the -5 eV to 190 eV energy range (0 eV is defined at 7130 eV). In the refinement, all atoms belonging to the histidine rings were moved rigidly linked to the N atom coordinated to the Fe. In the first step of the optimization process, only the bond distances were allowed to float. In subsequent steps the dihedral angles were also allowed to float. After each step of structural parameter refinement, a non-structural parameter refinement was performed. Many applications of MXAN to small molecule XAS data indicate that the correlation between structural and non-structural parameters is small. Nevertheless, the refinements were closely monitored for any large fluctuations in the structural and non-structural parameters during the refinement process. No large deviations in either the structural or the non-structural parameters were observed. The least-square error and a visual comparison of data and fit were used to determine the goodness of fit.

#### 2.5 EXAFS Data Analysis

Theoretical EXAFS signals  $\chi(k)$  were calculated using *FEFF* (version 7.0)<sup>23,24</sup> and the crystal structure of wild-type LO as initial model, and were fit to the data using EXAFSPAK (G. N. George, SSRL).<sup>21</sup> Theoretical paths corresponding to Fe-S and Fe-O distances of 2.1–3.0 Å (in 0.1 Å steps) were also calculated to simulate contributions of the S(Cys) to the EXAFS

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of wild-type LO and a new set of theoretical  $\chi(k)$  were calculated using *FEFF* and the data refit using the new parameters. The structural parameters varied during the fitting process were the bond distance (R) and the bond variance  $\sigma^2$ , which is related to the Debye-Waller factor resulting from thermal motion, and static disorder. The non-structural parameter E<sub>0</sub> (the energy at which k = 0) was also allowed to vary but was restricted to a common value for every component in a given fit. Coordination numbers were systematically varied in the course of the fit but were fixed within a given fit.

#### 3. Results and Analysis

#### 3.1 Fe K Pre-Edge

The Fe K pre-edge results from a quadrupole-allowed dipole-forbidden  $1s \rightarrow 3d$  transition. These transitions are usually very weak (~100 times weaker than the edge) but can gain intensity when the coordination deviates from centrosymmetry and gains dipole-allowed character due to Fe 4p mixing into the Fe 3d orbitals, as  $1s \rightarrow 4p$  transitions are electric-dipole allowed.<sup>25</sup> It has been shown by extensive studies on small molecule model complexes that the energy position and intensity pattern of the pre-edge transitions are signatures of the oxidation state and ligand coordination, respectively.<sup>25–27</sup> Ferrous 6C (Oh) complexes typically have a total integrated area of ~4 units over the pre-edge transition range. In 4C and 5C complexes the center of inversion is absent, leading to more intense pre-edge transitions, typically by a factor of 3–4. 4C and 5C complexes can be differentiated on the basis of the intensity distribution of the pre-edge transitions.

The Fe K pre-edge spectrum of ferrous N694C LO is compared to that of ferrous PAH<sup>T</sup> (wildtype phenylalanine hydroxylase (PAH)) and PAH<sup>R</sup> (substrate and cofactor bound PAH) in Figure 1 (Inset).<sup>28</sup> It has been shown using X-ray absorption, circular dichroism (CD) and magnetic circular dichroism (MCD) studies that the Fe center in PAH<sup>T</sup> and PAH<sup>R</sup> are 6C and 5C, respectively.<sup>29</sup> The total integrated area under the pre-edge transitions of N694C LO, PAH<sup>T</sup> and PAH<sup>R</sup> are 11.8(0.8), 8.1(0.4) and 13.9(0.8), respectively.<sup>30</sup> The individual peak energy and intensity of N694C LO, PAH<sup>T</sup> and PAH<sup>R</sup> are listed in Table 1. The total integrated intensity of N694C LO is similar to that of PAH<sup>R</sup> indicating that the active site Fe is 5C. These results are consistent with MCD data for N694C LO, which also indicate that the active site in N694C LO is consistent with a 5C ligand field.<sup>18</sup> However, the intensity ratio of the two preedge features is 1.44, which is closer to a 6C complex than a 5C complex (the intensity ratio of the pre-edge features in PAH<sup>R</sup> and PAH<sup>T</sup> are 1.61 and 1.39, respectively, see Figure S1). In strictly 5C  $C_{4v}$  complexes, the lower energy pre-edge feature associated with an Fe 1s $\rightarrow$  $d_z^2$  (A<sub>1</sub> symmetry) transition, gains intensity due to Fe  $4p_z$  (A<sub>1</sub> symmetry) mixing with the  $d_z^2$  orbital. However, in protein environments the active site can deviate considerably from an ideal  $C_{4\nu}$  geometry due to differences in ligands or the presence of a weak sixth ligand.<sup>31</sup> This lowers the symmetry and the higher energy pre-edge feature can also gain intensity through Fe  $4p_{x,y}$  mixing with the  $d_x^2 - v^2$  orbital. This indicates that the active site in N694C LO is more distorted than that in PAH<sup>R</sup>

Figure 1 shows a comparison of the rising edge region of N694C LO, PAH<sup>T</sup> and PAH<sup>R</sup>. The intensity of the peak at ~7126 eV is lower and similar for N694C LO and PAH<sup>R</sup> compared to PAH<sup>T</sup>. Since the intensity of this peak is characteristic of the number and type of first shell ligands,<sup>32</sup> the edge data of N694C LO indicates a ligand field consistent with a 5C structure and with the pre-edge data and MCD studies.

#### 3.2 EXAFS Region

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The EXAFS data for N694C LO was Fourier transformed over  $k = 2-12.5 \text{ Å}^{-1}$ . The EXAFS and the Fourier transform data are compared with that of the wild type (WT) sLO in Figure  $2.^{33}$  The EXAFS for WT sLO has a phase-shift to lower *k* relative N694C sLO similar to the relative phase shift in PAH<sup>T</sup> (6C) and PAH<sup>R</sup> (5C).<sup>29</sup> The first shell Fourier transform intensity has decreased by almost a factor of two in N694C LO relative to WT sLO. Both the WT sLO and N694C sLO data were collected in the presence of ~30% glycerol as the glassing agent. In the case of WT sLO, it has been shown using CD, MCD and K-pre-edge studies that in the presence of glycerol (or alcohols) the active site structure shifts from a 5C/6C (40%/60%) mixture to a purely 6C form. First shell Fourier-filtered EXAFS fits were also consistent with these results indicating 6 Fe-O/N contributions at ~2.16 Å<sup>17</sup> The EXAFS fits have been refined here using a full multiple scattering approach. The best fit to the WT-LO EXAFS and the corresponding Fourier transforms are presented in Figure S2, with the fit parameters given in Table S1. The first shell is fit using 6 Fe-O/N contributions at 2.18 Å.

In N694C LO, asparigine 694 is replaced by a cysteine residue. Thus, the weakly coordinating amide oxygen is replaced by a thiolate S(Cys) group which can potentially coordinate strongly with the Fe atom. However, due to the smaller size of S(Cys) relative to O(Asn), its approach to the ferrous center is restricted by the backbone in N694C sLO, compared to the more flexible approach of the amide O of asparigine in the WT sLO. This might create a possible pocket for a H<sub>2</sub>O molecule in N694C sLO which can coordinate to the Fe atom as a sixth ligand.<sup>34</sup> Both these possibilities were explored using appropriate models to generate theoretical EXAFS signals  $\chi(k)$  using FEFF. The fits to the EXAFS data using both these models and the corresponding Fourier transforms are shown in Figure 3A and 3B. The best-fit parameters are presented in Table 2. The Fe/S model was generated by including an Fe-S contribution at 2.7 Å in the input structure.<sup>35</sup> Using this model, the first shell was fit using 1 Fe-O/N path at 1.97 Å, 4 Fe-O/N paths at 2.12 Å and 1 Fe-S at 2.70 Å.<sup>36</sup> The second and third shells were fit using single and multiple scattering contributions arising from the histidine ring N and C atoms. The  $\sigma^2$  value for the Fe-S path (Table 2) was high indicating that, if present, the long Fe-S contribution is disordered. Fits using a coordination number of less than 1 for the Fe-S path improved the  $\sigma^2$  value but did not affect the goodness of the fit, consistent with a putative disordered Fe-S bond. For the fits to the Fe/O model, which was generated using a structure that included an Fe-O path at 2.5 Å, the best fit was consistent with 1 Fe-O path at 1.96 Å, 4 Fe-O/N paths at 2.12 Å and 1 Fe-O path at 2.49 Å. The second and third shells were fit using single and multiple scattering contributions arising from the histidine ring N and C atoms. Good fits were obtained when the third shell was been modeled with two sets of 8 Fe-N/C-N/ C multiple scattering components (adding the single scattering path did not improve the fit). However, the correlation between the bond distances and the  $\sigma^2$ -factors for these two multiple scattering paths were quite high, which might lead to the unusually low  $\sigma^2$ -factor value for the multiple scattering path at 4.28/4.29 Å. The correlation matrix obtained from the EXAFS fits to both the Fe/O and Fe/S model are included in the supporting information (bond distance= correlation between parameters 32 and 37 and sigma values= correlation parameters 33 and 38).

The  $\sigma^2$  value of the Fe-O path at 2.5 Å is moderately high which also indicates that, if the O is present, the Fe-O bond is disordered. The goodness of fit was very similar for both the Fe/S and Fe/O models. Thus, the EXAFS study indicates that the first shell is composed of 1+4 Fe-N/O paths and a weakly coordinated Fe-O or Fe-S path, which is disordered. This is consistent with the observed intensity distribution and the total area for the pre-edge transitions, since a five-coordinate site with a weak sixth interaction would have a symmetry lower than  $O_h$  and would gain allowed character from 4p mixing into the 3d manifold, which would

increase the intensity; and the presence of a weak interaction would change the intensity distribution from that observed for a pure  $C_{4\nu}$  species.

Thus, the EXAFS analysis indicates that the structure is distorted 6C with a weak sixth ligand, which can either be a distant Fe-O or Fe-S interaction. The Fe-S interaction would result from the S(Cys694) while the Fe-O interaction could result from two possibilities: *i*) a water molecule or, *ii*) reorientation of the ligated carboxylate group of the isoleucine to bind in an asymmetric bidentate fashion. Since the angular information available from EXAFS analysis is limited, both structural models for the Fe/O case are possible.

#### 3.3 Near-Edge Region

To investigate the presence and nature of the weak sixth ligand, near-edge simulations were performed using MXAN, which uses a multiple scattering approach to simulate the edge region.  $^4$  The N694C LO data were fit to 200 eV above the edge. Fits were first performed using the 5C active site structure obtained from the WT crystal structure. An iterative fit process of structural and non-structural parameters was performed, with the goodness-of-fit assessed as the square residual error  $(R_{sq})$ .<sup>8</sup> The  $R_{sq}$  for the 5C fit was 1.2. Figure S3 shows the best fit using the 5C structure, with the structural parameters given in Table S2 (Supporting Information). Fits were then performed by using both the Fe/S and Fe/O models (see the EXAFS section). For the Fe/S model, the simulations were very sensitive to the inclusion of an Fe-S component between 2.3-3.0 Å and the resultant fits were considerably worse (Figure 4A) with an R<sub>sa</sub> value of 3.9. Compared to the XAS spectra of a metal with O/N only ligands, there is usually a dramatic change in the XAS spectra of metal-S containing molecules, both in the rising edge and the near-edge region which is usually associated with an increase in covalent interaction between the metal and sulfur. Since the near-edge fit results cannot accommodate an Fe-S interaction, it provides a clear indication that Cys694 is not ligated to Fe in the resting ferrous state. Using the Fe/O model, two input structures were generated. In the WT crystal structure the carboxylate group (Ile) is asymmetrically coordinated with 1 Fe-O at 2.28 Å and the second Fe-O at 3.49 Å. The loss of the Asn ligand in N694C sLO might lead to a perturbation in the two Fe-O(Ile) bond distances, which can become shorter to compensate for the loss of the axial asparigine while still retaining their asymmetric coordination. Thus, the first input structure generated has two carboxylate Fe-O distances at 2.1 Å and 2.5 Å consistent with the two Fe-O bond distances obtained from the EXAFS data analysis. The second input structure consisted of a 6C structure with an axial Fe-O interaction at 2.5 Å representing an additional water molecule. Near-edge simulations using the asymmetric bidentate carboxylate structure resulted in an  $R_{sq}$  value of 3.7 (Figure 4B). This indicates that the carboxylate is coordinated in a monodentate fashion to the ferrous center with a sixth weak water ligand. Since the 6C fit with an asymmetric isoleucine carboxylate is worse than the 5C fit, near-edge simulations were performed with a sixth axial interaction at 2.5 Å (the equatorial plane is described by the 2 N(His), 1 O(Glu) and 1 O(H<sub>2</sub>O)). It has been shown that such axial M-O bonds are sensitive to dihedral angle made by the M-O bond with the equatorial plane.<sup>8</sup> Thus, the structure obtained from the 5C best fit was modified to obtain various 6C structures which included an Fe-O component at 2.5 Å and made different dihedral angles with the equatorial plane (see Figure S4, Supporting Information). These structures were then used to obtain structural and non-structural fits to the near-edge data. Figure 5 shows the data and best-fit spectrum. The inset shows the optimized structure, and the structural parameters are presented in Table 2.37 The best fit corresponds to an axial Fe-O distance of 2.50 Å, which is in good agreement with the EXAFS data. All the other first shell bond distances are in reasonable agreement with the EXAFS result. The R<sub>sa</sub> value is 0.53, which is considerably better than that for the bidentate carboxylate, and the Fe/S model strongly indicating the presence of a weak long Fe-O coordination. It should be noted that since the near-edge calculations are performed within the full multiple scattering approach, i.e., since

the inverse of the scattering path operator was computed exactly, (avoiding any 'a-priori' selection of the relevant multiple scattering paths) the individual contribution of the histidine multiple scattering paths cannot be obtained. However, as seen from the insets in Figure 3 and 4 (and cartesian coordinates given in supporting information), the orientation of the histidine rings remain reasonably similar in fits to the Fe/O and Fe/S models indicating that the increase in error value on going from the Fe/O to the Fe/S model results predominantly due to the replacement of the Fe-O with the Fe-S contribution. It is interesting to note that fits to both the Fe/O and Fe/S models are reasonably good in the low R region in the EXAFS analysis but differ significantly in the near-edge analysis. This is accounted for by the following factors: a) It has been previously observed that an Fe-S contributions affect the edge significantly due to considerable changes in the electronic structure. This is exploited by near-edge fitting routine implemented in MXAN, which calculates the inverse of the scattering path operator, exactly and treats the low energy region better; b) the Debye-Waller damping factor is not applicable to the near-edge fitting method using MXAN, thus, a "forced adjustment" of a longer Fe-S bond is not possible using a large sigma value and c) the EXAFS data are only available to k =12.5  $\text{\AA}^{-1}$  (typical for dilute non-heme iron proteins) which restricts our ability to distinguish between theoretically generated models with small differences.

Fits to EXAFS data usually give a 25% error in coordination number due to the strong correlation between the  $\sigma^2$  value and the bond distance. In addition, partial coordination (presence of a mixture of two components) can be explored using non-integer coordination numbers in the EXAFS fits. However, an exact two-component fit is currently unavailable in near-edge simulations using MXAN. To explore a 5C/6C mixture in N694C LO using near-edge simulations, a linear combination of the best fit 5+1C and the 5C model was performed within a window of 10%6C/90%5C and 90%6C/10%5C. These simulations were then

compared to the data and the error (e) was obtained as  $\varepsilon = \sum_{0-200eV} |S_{exp} - S_{sim}|$ .

Figure S5 (black line) (Supporting Information) shows the change in error value with an increase in the 5C component. A corresponding EXAFS fit with non-integer Fe-O coordination was also performed and the change in error value with an increase in 5C component is shown in Figure S5 (red line). These results clearly indicate that the molecule is 5+1C with no observable pure 5C component. Interestingly, the error for 10% to 50% 5C component (higher amount of 5C component leads to unreasonable  $\sigma^2$  values) remains very similar for the EXAFS fits while that for MXAN shows a clear decrease in the goodness of fit. This indicates that edge-fitting is more sensitive to partial coordination number. Thus, these results emphasize the sensitivity of the multiple scattering method in fitting the near-edge region and obtaining geometric structure information, in particular angular information, which is usually inaccessible using EXAFS.

#### 4. Discussion

#### 4.1 Geometric Structure of N694C sLO1

In this study, a combination of pre-edge, near-edge and EXAFS studies have been performed on N694C sLO to determine the active site geometric structure. These results indicate that the ferrous center is 5+1C with a weak Fe-O(H<sub>2</sub>O) coordination at 2.5 Å. It is interesting to note that the MCD results indicate that the site is 5C with two ligand field bands at 6050 cm<sup>-1</sup> and 11450 cm<sup>-1</sup>, although the MCD studies indicate a small increase in the ligand-field on going from WT to N694C sLO. This discrepancy in the MCD and XAS results stem from the fact that a long Fe-O distance at 2.5 Å does not affect the bonding significantly enough to be seen as a coordination number change in MCD. However the small shift in the MCD bands to higher energy indicates an increase in ligand field, reflecting a presence of a weak sixth ligand.

Both the WT and N694C sLO EXAFS data were measured in the presence of ~30% glycerol. It has been shown in previous studies that glycerol converts the resting site (5C) to 6C, which is similar to the structural change observed upon substrate binding.<sup>17</sup> This indicates that mutation of the weakly coordinated asparigine ligand to the smaller S(Cys) ligand removes the ability of the protein to convert into a 6C structure upon substrate binding. This is important in the reactivity of the protein since this 5C to 6C conversion, which is attributed to the coordination flexibility in the site, plays a key role in the catalytic rate. The weakly bound  $H_2O$  molecule keeps the site 5+1C in both resting and substrate bound forms in N694C sLO1. The presence of this H<sub>2</sub>O molecule likely modifies the H-bonding network between the second sphere residues by introducing a H-bond between E697 and the H<sub>2</sub>O molecule.<sup>38</sup> This H-bond replaces the one lost between E697 and N694 due to mutation. Thus, although the substrate docking associated with the H-bond network might not be perturbed significantly, the substrate bound active site structure (simulated by the presence of glycerol) changes dramatically between WT and N694C sLO as observed from a comparison of the EXAFS data (see Figure 2). This leads to the observed ~3000 fold decrease in the catalytic rate in N694C lipoxygenase relative to WT sLO.<sup>18</sup>

#### 4.2 Comparison of Edge and EXAFS Analysis

The results presented in this study show that near-edge analysis can be used to complement EXAFS results and even give structural information inaccessible by EXAFS analysis. In N694C lipoxygenase the two six-coordinate models involving a sixth oxygen atom, an axial water or a bidentate carboxylate of the Ile residue, can only be evaluated using a full multiple scattering approach, which is capable of determining three-dimensional structural information. At this point it is important to evaluate the maximum number of independent parameters ( $I_P$ ) and the reliability of the fit for the EXAFS and near-edge MXAN analysis. In the EXAFS

region I<sub>P</sub> can be directly calculated using the relation  $I_p = \frac{2\Delta k\Delta R}{\pi} + 2$ . <sup>39</sup> For the N694C lipoxygenase EXAFS data I<sub>P</sub> is ~20. The I<sub>p</sub> number used in the EXAFS analysis presented here is 15, which is well within the upper-limit of I<sub>p</sub> that can be used to obtain an acceptable EXAFS fit.

In the edge region, however, since the mean free path of the photoelectron is very large, the multiple scattering contributions can extend up to  $\sim 10$  Å. This can exponentially increase the number of possible single- and multiple-scattering contributions to the edge region. Inclusion of all these paths can lead to structure-optimization problems since the possibility of reaching a local minimum for a system with a large number of independent parameters also increases rapidly. In the near-edge analysis presented here, single and multiple scattering contributions up to 4.5 Å have been considered, using 18 structural and 4 non-structural parameters, respectively. It has been shown that the structural and non-structural parameters have very low correlation, thus, the number of independent structural parameters in this analysis is ~18. To justify the use of atoms up to 4.5 Å from the central ferrous atom, a shell-by-shell analysis is presented in Figure 6, which shows multiple scattering contribution up to 2.6 Å (Figure 6A), 3.5 Å (Figure 6B) and 4.5 Å (Figure 6C, final fit). For each shell, the structural parameters were allowed to float. The 2.6 Å fit is very poor with an  $R_{sq}$  of 6.5. While the 3.5 Å model reasonably fits the experimental spectral features ( $R_{sq}$  =1.7), the final 4.5 Å fit range ( $R_{sq}$ =0.53) is essential to obtain the double-peak feature between 50–80 eV. Thus, the results  $\frac{1}{2}$ indicate that multiple-scattering contributions up to 4.5 Å are essential to successfully model the rising-edge and near-edge region. This is in agreement with the EXAFS data, which require multiple-scattering contribution from the  $C_{\beta}$  and  $N_{\beta}$  in the 4–4.5 Å range for a good fit. However, it is not sufficient to show that multiple-scattering contributions from atoms at a given distance are important to fit the edge-data. Structural parameters in a fit can be highly correlated leading to fits that do not represent the global minimum. The correlation matrix

between the free parameters can be obtained using the MINUIT code implemented in the MXAN package and that for the best-fit shown in Figure 6C and Figure 5 is presented in the Supporting Information. The matrix indicates low correlation between most parameters and supports the fact that the structural fit obtained using MXAN is physically justifiable. It is important to note that examination of the correlation-matrix is important in EXAFS fitting procedures also, and can give an estimate of the softness of the best-fit structural parameters. Often, a particular bond distance in the EXAFS fit is closely correlated with its corresponding  $\sigma^2$  value. Since the a-priori estimation of  $\sigma^2$ s for individual systems is not yet possible, an acceptable range of values is available for any particular absorber-scatterer pair based on EXAFS analysis on small models. However, this range of acceptable  $\sigma^2$  values is quite large and, in some cases can lead to large errors in bond-distance estimation, especially in the case of second and third shell single- and multiple-scattering. In most cases, a combination of chemical knowledge and close inspection of the statistical parameters at every step of the fitting process is important for accurate structure determination.

In summary, this study describes the application of multiple-scattering edge-fitting methodology to a non-heme ferrous active site of the N694C mutant of lipoxygenase. It is shown that edge-fitting analysis can give important three-dimensional structural information, which usually cannot be accessed using EXAFS analysis. In particular, for systems which are subject to limitations that can severely affect the resolution and accuracy of EXAFS analysis. In addition, since the  $\sigma^2$  damping terms in the XANES energy region are almost temperature-independent, atomic vibrations have negligible effects and do not interfere with structure determination. This allows precise determination of partial coordination numbers and can be helpful in determining different compositions in mixtures.

#### **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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- 36. A strong Fe-S(Cys) bond could lead to weakening of the H<sub>2</sub>O ligand bond, leading to a distorted 5C structure with 4 Fe-O/N, 1 Fe-S and 1 long Fe-O interaction. However, EXAFS data could not be reasonably fit using a structural model reflecting this geometry.
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#### Figure 1.

The normalized Fe K near-edge region of N694C LO (—),  $PAH^R$  ( \_) and  $PAH^T$  ( \_). Inset shows the expanded pre-edge region.





Fourier transforms (non phase shift corrected) and EXAFS data (Inset). N694C LO (—), WT-LO ( \_).



#### Figure 3.

Fourier transforms (non phase shift corrected) and EXAFS data (Inset) for N694C LO. A: Best fit using the Fe/S model. B: Best fit using the Fe/O model (see text for more details). Data (—), Fit ( \_\_).

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#### Figure 4.

The normalized Fe K near-edge spectrum of N694C LO (—) and the best fit ( \_) obtained from the optimization of both structural and non-structural parameters. (A) Fe/S fit. (B) Fe/O bidentate carboxylate fit.





The normalized Fe K near-edge spectrum of N694C LO (—) and the best fit (  $\_$ ) obtained from the optimization of both structural and non-structural parameters using the Fe/O axial water model.





#### Figure 6.

The normalized Fe K near-edge spectrum of N694C LO ( \_\_) and the best fit (\_\_) obtained from the optimization of both structural and non-structural parameters using the best-fit model (six-coordinate Fe<sup>II</sup> with axial water) up to 2.6 Å (A), 3.5 Å (B) and 4.5 Å (C).

#### Fe-K Pre-edge Analysis.

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	N694C-LO	PAH <sup>R</sup>	PAH <sup>Tb</sup>
Peak 1 (eV)	7111.7	7111.7	7111.8
Peak 2 (eV)	7113.5	7113.5	7113.7
Total Area <sup>C</sup>	11.8(0.4)	13.9(0.8)	8.1(0.4)
Pre-edge Ratio	1.44	1.61	1.39
Edge maxima (eV)	7126.3	7126.4	7125.7

 $^{a}$  Values in parentheses are the statistical standard deviations calculated from the individual acceptable fits used in the analysis.

 $^{b}$ The first two peaks in PAH<sup>T</sup> are in the limit of resolution and hence treated as a single peak for relative ratio estimations.

<sup>c</sup>The reported values are multiplied by 100 for convenience.

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 Table 2

 EXAFS Least Squares Fitting Results for N694C Lipoxygenase.

F1(#	Coordination/Path	$R(\hat{A})^d$	$s^2 (\dot{\Lambda}^2) b$	E <sub>0</sub> (eV)	$F^{C}$
	1 Fe-N/O	1.97(0.02)	259(48)		
	4 Fe-N/O	2.12(0.02)	442(202)		
	1 Fe-S	2.70(0.03)	1300(131)		
Model Fe/S	6 Fe-C	3.12(0.02)	557(44)	-6.99	0.135
	14 Fe-N-C	3.27(0.02)	557 <sup>d</sup> (44)		
	8 Fe-N/C-N/C	4.29(0.04)	163(190)		
	8 Fe-N/C-N/C	4.44(0.03)	719(190)		
	1 Fe-N/O	1.96(0.03)	420(113)		
	4 Fe-N/O	2.12(0.02)	520(66)		
	1 Fe-O/N	2.49(0.02)	542(121)		
Model Fe/O	6 Fe-C	3.11(0.02)	725(65)	-7.99	0.138
	14 Fe-N-C	3.29(0.04)	725(65)		
	8 Fe-N/C-N/C	4.28(0.04)	279(240)		
	8 Fe-N/C-N/C	4.41(0.02)	855(240)		
<sup>a</sup> The estimated standard deviations for	r the distances and Debye-Waller calculated by E	XAFSPAK are shown in paranthesis.			

 $^{b}$ The  $\sigma^{2}$  values are multiplied by  $10^{5}$ .

<sup>c</sup> Error is given by  $\Sigma[(\chi_{obsd}-\chi_{calcd})^2 k_6]/\Sigma[(\chi_{obsd})^2 k_6]$ .

 $^{d}$ The  $\sigma^{2}$  factor of the multiple scattering path is linked to that of the corresponding single scattering path. The S $_{0}^{2}$  value was fixed at 1 for all refinements.

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Table 3

The estimated standard deviation for the distances is  $\pm 0.01$  Å and bond angles (not shown) is 4°.

a'The structural and non-structural parameters are obtained by alternate fitting of each set pf parameters separately. This is possible due to the low correlation between these parameters. Nax=axial N (His), Neq=equatorial N(His).

<sup>b</sup>Fe-OCO<sup>-</sup>(Ile).