Biophysical Characterization of a Disabled Double Mutant of Soybean Lipoxygenase: The “Undoing” of Precise Substrate Positioning Relative to Metal Cofactor and an Identified Dynamical Network

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ABSTRACT: Soybean lipoxygenase (SLO) has served as a prototype for understanding the molecular origin of enzymatic rate accelerations. The double mutant (DM) L546A/L754A is considered a dramatic outlier, due to the unprecedented size and near temperature-independence of its primary kinetic isotope effect, low catalytic efficiency, and elevated enthalpy of activation. To uncover the physical basis of these features, we herein apply three structural probes: hydrogen-deuterium exchange mass spectrometry, room-temperature X-ray crystallography and EPR spectroscopy on four SLO variants (wild-type (WT) enzyme, DM, and the two parental single mutants, L546A and L754A). DM is found to incorporate features of each parent, with the perturbation at position 546 predominantly influencing thermally activated motions that connect the active site to a protein-solvent interface, while mutation at position 754 disrupts the ligand field and solvation near the cofactor iron. However, the expanded active site in DM leads to more active site water molecules and their associated hydrogen bond network, and the individual features from L546A and L754A alone cannot explain the aggregate kinetic properties for DM. Using recently published QM/MM-derived ground-state SLO-substrate complexes for WT and DM, together with the thorough structural analyses presented herein, we propose that the impairment of DM is the combined result of a repositioning of the reactive carbon of linoleic acid substrate with regard to both the iron cofactor and a catalytically linked dynamic region of protein.

INTRODUCTION

Enzymes catalyze chemical reactions with enormous rate enhancements nearing $10^{26}$-fold that are accompanied by high regio- and stereospecificities relative to their uncatalyzed counterparts. The physical bases for such properties are of considerable ongoing interest and investigation. De novo designed protein catalysts, based on structural information of active sites and the principles of transition state theory, have generally achieved only moderate rate accelerations in the range of $10^2$–$10^6$-fold, even after further functional optimization via directed evolution. The large gap between naturally occurring catalysts and designed systems underscores the need for increased understanding that can be used to inform greater success in designed catalysts. Soybean lipooxygenase (SLO) is one of several paradigmatic systems for studying the properties of enzymatic room-temperature hydrogen tunneling and their relevance to general, multi-dimensional models for the origins of enzymatic rate acceleration. SLO catalyzes the stereospecific oxidation of linoleic acid through a rate-limiting abstraction of a pro-S hydrogen atom from the C11 of the substrate linoleic acid (LA); this occurs via a proton coupled electron transfer (PCET) reaction in which a proton is transferred to the active site ferric-bound hydroxide and an electron reduces the ferric center to ferrous (Scheme 1). The initially measured and,
The temperature-dependence of the KIE, as assessed as I553G, showing an increase in the KIE to ca. 180°. A generating a enthalpy of activation for D-transfer that can deep-tunneling kinetic parameter correlations to be made between active site motions and the experimental KIEs during the biomolecular reaction of enzyme—substrate (E-S) substates. For native enzyme, one of the primary functions of conformational sampling is the transient achievement of hydrogen donor and acceptor distances that are sufficiently reduced from those of initial van der Waals complexes to enable productive wave function overlap.

Studies of a series of double mutants of SLO have further pushed the envelope in the ability of theoretical models to rationalize behavior.22–29 While the majority of double mutants show behavior similar to single mutants, i.e., analogous KIE values at RT that are accompanied by an increase in ΔEa (Table S1), a single dramatic outlier, L546A/L754A (DM), emerged with a kfi/kfo of 537 ± 55 under presteady single turnover anaerobic measurements29 and an average kfi/kfo of 661 ± 27 via steady-state measurement across six temperatures.28 Additionally, in contrast to the KIEs of other active site single and double mutants that show temperature sensitivity, the KIEs in DM change little if at all with temperature (Table S1). These properties of DM SLO can be well-reproduced using the developed vibronically nonadiabatic PCET model, where donor—acceptor distance sampling within the active site appears to have become rigidified compared to WT and the other SLO variants,22,28,29 preventing DM from recovering the tunneling-ready distance characteristic of native enzyme and other well characterized mutants. A related temperature—pressure perturbation study of DM SLO also highlights the distinctive features of this variant.26 Unlike the pressure-induced shifted conformational landscape in WT and three single mutants (L546A, L754A and I553V), the conformational landscape of the enzyme—substrate complex of DM is more resistant to perturbations from elevated pressure, showing a consistently high energy barrier (Ea = 8—9.9 kcal/mol) and slow catalytic efficiency (104-fold decrease) relative to WT. At this juncture, a major unanswered question is the structural origin underlying the aggregated features of DM SLO.

In this study we compare a suite of biophysical probes applied to DM, the two components of DM (L546A and L754A) and WT. These probes include HDXMS as a function of temperature, EPR spectroscopy and room temperature X-ray crystallography. The primary results show that DM adopts features from both parent single mutants, exhibiting the same HDXMS behavior as L546A and showing structural changes similar to L754A. Significantly, in no instance has evidence been found for enhanced structural rigidification in the resting form of enzyme, in contrast to the earlier inference of rigidification within the donor—acceptor distance sampling modes. As we discuss, the aggregate data indicate a disruption in the positioning of the reactive carbon of substrate with regard to the catalytic iron center that is further incapacitated via an impairment of its communication with a dynamical network that has been identified to extend from the active site to a remote protein/solvent interface.
RESULTS

Hydrogen–Deuterium Exchange. The time and temperature dependent analysis of HDXMS was carried out with DM for comparison to the previously characterized WT SLO.\(^{33}\) Using the same set of nonoverlapping peptides selected for WT SLO, 78% of the protein sequence for the DM catalytic domain (C-terminal domain) was covered (compared to 89% for WT). The apparent averaged rate constants for exchange, \(k_{\text{HDX(avg)}}\), from exponential fits to the HDXMS traces for WT and DM are presented in Table S2 and represent the protein in the absence of substrate and exchanged in buffer pH L (L = H, D) of 8, conditions where EX-2 conditions have been demonstrated. EX-2 conditions enable us to extract thermodynamic information regarding regional peptide flexibility. The majority of DM-derived peptides display nearly identical exchange behavior to WT across the 5 temperatures (10, 20, 25, 30, and 40 °C) analyzed (cf. Figure S1). In a previous study, peptide 317–334, a solvent-exposed loop (Figure 1A), was shown to produce significant alterations in \(E_a\) for \(k_{\text{HDX(avg)}}\) following mutation at positions 553 (I553G) and 546 (L546A).\(^{33}\) This peptide exhibited trends in the \(E_a\) for hydrophobic and catalytic PCET (solid line in Figure 1C).\(^{33}\) In the current HDX analysis of peptide 317–334 in DM, we observe a value for \(E_{\text{HDX(avg)}}\) again distinct from WT (Figure 1B, Table S3); however, while DM SLO displays the highest enthalpic barrier of catalysis among all SLO variants (Table S1), the value of \(E_{\text{HDX(avg)}}\) associated with DM is 14 kcal/mol, almost identical to the behavior of L546A (Figure 1C, top red arrow). This initially surprising result, of comparable \(E_{\text{HDX(avg)}}\) values in DM and L546A, implies that the added active site packing defect that arises from L754A in DM does not impact the thermally activated surface motions that correlate with catalysis.

To further substantiate the lack of an impact of L754 on the long-range protein motional network, the single variant L754A was examined with regard to the time and temperature dependence of its HDXMS. The L754A demonstrates an elevated enthalpic barrier of catalysis (\(E_{\text{HDX(avg)}} = 4.1\) kcal/mol) and an even more impaired catalytic rate constant than L546A (Table S1);\(^{21}\) however, the \(k_{\text{HDX(avg)}}\) and \(E_{\text{HDX(avg)}}\) obtained from the HDXMS profiles of all L754A derived peptide across the full temperature range are within experimental error of WT (Table S2, S3) (Figure 1C, bottom red arrow). The absence of a linkage between the \(E_{\text{HDX(avg)}}\) and \(E_{\text{HDX(avg)}}\) in L754A supports an uncoupling between the packing defect at position 754 and motions at the protein–solvent interface, such that the observed rate impairment with this mutant must arise from other features (see below). We note that there were no noticeable changes in the extent of exchange as a function of temperature for L546A, L754A and DM (cf. full data sets in Figure S1), in marked contrast to the previously studied I553X series where the primary impact of mutation was seen as an increase in the extent of exchange in the S41–S65 region of protein.\(^{33}\)

These new comparisons between HDXMS profiles and catalytic parameters highlight the combinative yet distinct contributions from the residues L546 and L754 in defining the behavior of DM. As earlier noted, L546 is located within the identified network of structural communication between the surface loop (317–334) and active site residues (Figure S2),\(^{33}\) with a packing defect at position 546 influencing motions at the protein–solvent interface that are reflected both in \(E_{\text{HDX(avg)}}\) and \(E_{\text{HDX(avg)}}\). This impact from L546A appears well preserved in DM. In contrast, L754 resides near the iron cofactor and spatially away from the identified network (Figure S2), suggesting that changes in the microenvironment around the metal, such as the degree of hydration, ligand field, and/or redox potential may be the major cause of the observed change in properties for L754A. These possibilities are explored below.

Room-Temperature X-ray Crystallography of the SLOs. To systematically reveal structural changes associated with the size-reduction at position 546 and/or position 754, room temperature X-ray crystallography was adopted for WT, L546A, L754A and DM. Such room temperature structures can reveal a multiplicity of protein conformers that are undetectable under cryo-conditions.\(^{41,42}\) Additionally, in the context of visualizing solvent water, difference maps calculated from two room temperature structures are more reliable than cryogenic maps.\(^{33}\) High resolution (1.70–1.85 Å) data were obtained for L546A, L754A and DM, revealing practically superimposable protein backbones and largely unaltered side-chain conformations in comparison to the room temperature X-ray structure of WT.\(^{33}\) The most dramatic change in these three mutants is the expanded active site that is associated with changes in solvation. The size reduction of the side chain on position 546 and/or position 754 leads to a gradually enlarged...
active site. Defining a more specific active site pocket than in the earlier study, the cavity in WT is calculated to be \( \leq 21 \text{ Å}^3 \). Using this new value as a frame of reference, the variants described in this study indicate increases of 170% in L546A, 76% in L754A, and over 270% in DM. The isomorphous difference map was further applied to individual mutants using phases from the room temperature WT model. The difference map between WT and DM indicates an increased electron density in the active site of DM (Figure S3A), resembling most closely the difference map between WT and L754A (Figure S3B). These density changes imply increased bound (or diffuse) water molecules near the iron cofactor. The other single mutant L546A demonstrates a slightly positive density, which also implicates the presence of extra water molecules (Figure S3C). However, the number of water molecules visualized is clearly less than for DM and L754A, and restricted to a position near 546 and H499. It is important to note that changes in solvation have not always been associated with mutation at hydrophobic side chains in SLOs: the I553X series failed to indicate X-ray evidence for the presence of any additional water molecules at low temperature and room temperature, albeit with a significantly enlarged volume near the mutated residue.26,33

To further substantiate possible microenvironment changes around the iron cofactor, we turned to a detailed inspection of the impact of mutation on the active-site geometries for WT, L546A, L754A and DM. Figure 2A–D indicates no significant changes in the coordination geometry between the iron and first/second sphere residues in all SLOs studied. The major difference within the active site is the newly entered water molecules in each variant that may interact with other coordinating residues through hydrogen bond networks. In both L546A and L754A, one additional water molecule is seen to occupy the packing defect created near position 546 and 754, respectively. However, the newly entered water molecule in L754A forms hydrogen bonds with two first sphere coordinating residues (iron-bound water molecule and H499) as well as a second sphere coordinating residue Q495, while the extra water in L546A is far from the second sphere coordinating residue Q495, interacting dominantly with H499 (Figure 2B,C). The water molecules occupying these single site packing defects and their associated hydrogen bond network are retained in DM. Beyond these two water molecules, we further observe two additional water molecules in DM that are adjacent to position 754 and form hydrogen bond network to the first sphere coordinating residues I839 as well as Q697.

**EPR Spectra of Fe(II) and Fe(III) SLOs.** In order to further explore changes in the microenvironment around the nonheme iron center of SLO that result from L546A, L754A and DM we employed EPR spectroscopy. The WT, L546A, L754A and DM isolated from *E. coli* all contain EPR-silent Fe(II), as shown in Figure S4, consistent with previous studies on SLOs and other lipooxygenases. The oxidized (“yellow”) Fe(III) SLOs were prepared through the addition of 2 equiv of substrate, linoleic acid (LA), to the Fe(II) SLOs as described in Materials and Methods section. The Fe(III) WT and L546A both demonstrate a complex multicomponent EPR signals with a pronounced feature around \( g = 7.3 \) and \( 6.2 \) and an additional weak feature at \( g = 4.3 \), Figure 3. The
previous EPR analysis of WT attributes the two groups of peaks to different geometric structures around the high-spin iron with the $g = 6.2$ signal representing axial asymmetry, $g \approx 4.3$ arising from a rhombic species, and $g = 7.3$ as an intermediate between the above two species. These features indicate that the ferric form of WT and L546A both contain the above-mentioned three species with axial asymmetry of the zero field splitting as the dominant feature.

Remarkably, once again the L754A shows distinct patterns from WT and L546A. The EPR spectrum of L754A shows dramatically increased intensity at the $g \approx 4.3$ peak and a weak feature at $g = 7.3$ (Figure 3), indicating that the ferric active site in L754A contains a primarily rhombic species. The interconversion between axial to rhombic species zero field splitting in lipoxygenases indicates the alteration in the active site geometry, which is usually seen to arise as the result of either product coordination to the iron center or substitution of the iron ligands that impacts the ligand field strengths. We propose that the unique EPR signals in L754A are directly related to the changes in microenvironment around the iron center, including the additional water molecule near the position 754 and its interaction with other iron ligands as shown in Figure 2C.

Unlike any previously reported high-spin ferric iron, the EPR studies of oxidized DM demonstrate a dramatically decreased EPR signal intensity (Figure 3). However, the increase in UV/vis Fe(III) absorption between 330 and 350 nm for DM, representing the formation of Fe(III) following incubation of enzyme with linoleic acid, appears almost identical to WT after correction for their respective iron levels (Figure S6); this indicates that the active site in DM SLO does contain an oxidized, Fe(III) species. The presence of Fe(III) in DM SLO is further validated by the strong peak at $g = 4.3$ after addition of 4-nitrocatechol to free enzyme (Figure 4D, top two EPR spectra). As shown previously in spectroscopic, kinetic and
structural analyses, 48–50 4-nitrocatechol can bind to the active site of SLO and form complexes with Fe(III) SLO in the absence of a redox reaction at pH = 7.0. The dramatically increased intensity in the 4-nitrocatechol complex is, thus, attributed to a zero field splitting change caused by displacement of first sphere side chain ligands with catechol oxygens within the DM-4-nitrocatechol complex. 48 As has been reported for SLO and homologues of lipoxigenase, the Fe(III) spectra can have a range of zero field splitting parameters (D = −0.3 to +1.7 MHz, E/D = 0.02 to 0.33) which is influenced by the microenvironment of the Fe and will greatly affect the intensity and shape of the EPR spectrum. 46,47 We note that the 10x EPR spectra of oxidized DM SLO displays features of both WT/L546A and L754A (Figure 3).

Estimation of Redox Potential in SLOs. Room temperature X-ray structures, together with the above EPR analyses, indicate significant disruption in both the microenvironment and electronic structures of the iron sites in L754A and DM. Such changes could impact the redox potential of the active site iron, which would further impact both the reaction free energy ΔG° and the corresponding rate constant kcat. A previously developed method by Nelson demonstrated that formation of complexes between oxidized WT Fe(III) SLO and a series of substituted catechols enables an estimate of the redox potentials of the active site iron. 50 Specifically, conditions were established for the SLO-catechol complexes, in which less reducing catechols do not transfer an electron to the active site Fe(III), while the more reducing catechols reduce the ferric iron to EPR silent Fe(II), establishing a redox ruler for the active site iron of SLO.

Figure 4A shows a series of EPR spectra of the WT SLO-catechol complexes after an incubation time of 12 h in phosphate buffer (pH = 7.0). Samples of complexes of WT with 4-nitrocatechol or 3,4-dihydroxybenzonitrile both show sharp EPR signals at g ≈ 4.3, which suggest a conversion from the dominant axial component of Fe(III) to a rhombic component of Fe(III) in the complexes. The EPR signals at g ≈ 4.3 are also seen in the complex of ferric WT SLO with 3,4-dihydroxybenzaldehyde albeit with largely decreased intensity, as evidence for a partially reduction of Fe(III) to Fe(II). In the case of the other five SLO-catechol complexes, the EPR spectra all demonstrate very minor peaks in the same region, indicating almost fully reduced Fe(II) in those complexes. The remaining very small peak have been suggested to arise from some portion of denatured protein that releases the free Fe(III). 51 The trends of the reaction between active site iron and catechols at pH = 7.0 are consistent with Nelson’s previous observations, and an assignment of a redox potential for iron in WT at pH = 7.0 between the range of the redox potentials for the test ligands (0.57 to 0.68 V). 50

We proceeded to measure the redox potential of iron in L546A, L754A and DM in a similar fashion, Figure 4B–D. In the cases of L546A and DM (Figure 4B,D), the trends in the EPR with 4-nitrocatechol and the more reducing catechol are similar to WT, implicating redox potentials for iron in these two mutants within the 0.57 to 0.68 V. The slight differences in the reducing ability of catechols in the middle range (Figure 4A,B,D, from 3,4-dihydroxybenzonitrile to caffeic acid) further imply a trend of redox potential that is DM > WT > L546A. Finally, in the case of L754A, the observation of a reduction of the ferric center by the least reducing reagent in this series, 4-nitrocatechol, indicates a potential that is somewhat above 0.68 V, for a full trend in redox potential of L754A > DM > WT > L546A.

Even though the adopted method can only give an estimated range and relative trend of redox potentials for SLO variants, we can conclude that the redox potential at the iron center does not directly control C–H activation rate constants. This is supported first, by only minor alternations in the range of redox potential (less than 0.1 V) for changes in rate constants of 108–109-fold. Second, the trend of the redox potential change (L754A > DM > WT > L546A) does not align with the trend of observed rate constants (WT > L546A > L754A > DM). Clearly, other factors dominate the enormous impact of the DM (and other variants) on the rate impairment of SLO.

### DISCUSSION

Hydrogen tunneling offers a unique window into the role of protein conformational motions in facilitating optimal active site alignment that accounts for the rate enhancements in enzyme catalyzed reactions. SLO has served as an ideal model for understanding the role of protein structure and dynamics in generating highly efficient tunneling rates and properties relative to small molecules in condensed phase. Kinetic analyses of this system over decades have led to a predictive multidimensional kinetic model that incorporates a hierarchy of protein motions. 5,9,16,35 Further, recent biophysical studies of WT versus single site SLO mutants, 33,34 have begun to reveal the molecular basis of catalysis-linked protein motions. The unique kinetic features emerging from a nearly fatal double mutant, DM, earlier led to the proposal of an elongation of the hydrogen transfer distance within its E-S complex that is accompanied by an apparent rigidification of the enzyme active site that prevents recovery to a tunneling DAD that approximates WT. These parameters include a greatly reduced kcat together with an enlarged temperature dependence on kcat (Ea ~ 10 kcal/mol) and an enormous KIE (kcat ~ 500–700) that is nearly independent of temperature (ΔEa ~ 0 kcal/mol). 28,29 The present work addresses the structural origin of the enormous KIE and other aberrant properties of DM SLO via the interrogation of a range of biophysical and structural probes that include HDXMS, EPR and room temperature X-ray crystallography.

Among the kinetic parameters characteristic of DM, kcat is found to be reduced 104-fold relative to WT. Previously, it had been shown that altering the ligands to the metal in SLO both reduced rates and decreased reduction potentials of the metal, implicating an altered chemical driving force, ΔG°, in catalytic rate impairment. 52 To test this possibility for the present series of SLO mutants, we examined relative reduction potentials using an established EPR method involving analysis of Fe(III) SLO complexes with various catechol derivatives. As shown, the DM undergoes only a very modest variation in its redox potential, with no systematic trend in the magnitude of its kcat and estimated E° relative to WT (Figure 4). These experiments serve as an important frame of reference in ruling out significant alterations in the chemical driving force as the source of the impaired activity of DM (see below).

**Origins of the Elevated Enthalpic Barriers in SLO Mutants.** Because wave function overlap from the rate-limiting C–H bond cleavage step in SLO catalysis is, by definition, barrier-less, the observed enthalpic barriers are derived from the conformational sampling of the enzyme–substrate complexes. We first examined the extent to which the aggregate behavior of DM could be rationalized from the two
parent mutants, L546A and L754A. Though these single mutants have the same enthalpic barriers ($E_{\text{hit}} = 4 \text{ kcal/mol}$), the origins of the increased $E_r$ values relative to the native enzyme ($E_{\text{hit}} = 2 \text{ kcal/mol}$) are now concluded to be different, arising either from impairment of a catalytically linked dynamic network in SLO (L546A$^{33}$) or an alteration in the active site microenvironment (L754A, this study).

HDXMS previously showed that reduction in side chain volume at L546 or I553 produces changes in the enthalpic barrier for HDXMS rate constants within an identical region of solvated protein ($317\text{–}334$) that correlate with trends in the enthalpic barrier for the catalytic PCET process.$^{33}$ The elevated catalytic $E_{\text{hit}}$ for L546A was, thus, assigned to an impact of this mutation on a catalytically linked protein dynamical network that traverses a distance of $\geq 20 \text{ Å}$ from the active site to the protein–solvent interface. By contrast, the current comparison of HDXMS profiles for L754A to WT shows these to be practically identical for every peptide including $317\text{–}334$, ruling out perturbations to the dynamical network as the primary origin of the altered properties of L754A.

The present studies point instead toward a perturbation of the microenvironment surrounding the iron cofactor as the source of changes in catalysis for L754A. The estimated relative redox potential for L754A (Figure 5) is seen to exhibit a modest increase relative to the $E^o$ of WT ($\Delta E^o < 100 \text{ mV}$), predicting an increase in rate, thereby making a change in redox potential an improbable source of its $10^3$ lower $k_{\text{cat}}$ and elevated $E_{\text{hit}}$. The position of the additional, structured water that infiltrates the expanded active site in L754A (Figure 2) may be a particularly significant feature that influences both the kinetic parameters and EPR spectrum of apo-L754A; the latter is clearly divergent from both WT and the L546A variant (Figure 3).

Buried water molecules have been assigned to important roles in biological function, including ligand binding, protein stability and flexibility.$^{55}$ In the case of buried waters that become displaced upon ligand binding,$^{58}$ these can provide an important thermodynamic driving force for enhanced substrate affinity (see for examples, refs 55–57, 59). WT SLO has at least five X-ray-resolved water molecules that line the putative ligand binding pocket$^{59}$ and most, if not all, are likely to be expelled upon substrate acquisition. Because of the unique positioning of the new water introduced upon mutation at position 754, it is possible that this will remain bound in the E-S complex, leading to changes in the active site dielectric that impact both $k_{\text{cat}}$ and the accompanying $E_r$. While X-ray crystallography of complexes of LA with SLO could, in principle, resolve any ambiguity regarding the degree to which active site waters are retained in the E-S complex (WT or otherwise), decades of efforts to obtain such data have thus far been unsuccessful. We note that X-ray structures of the apo-form of I553G provide a reasonable control for the impact of L754A. While the S53G variant shows an enlarged active site cavity volume similar to 754A, it lacks any detectable added water molecules by either cryo- or room temperature X-ray studies$^{25,33}$ and exhibits a turnover number much closer to WT.$^{26}$

The combination of impaired dynamics and altered active site hydration that are distinctive of L546A and L754A, respectively, begin to rationalize the greatly increased activation energy ($\sim 10 \text{ kcal/mol}$) and decreased rate of DM relative to WT. However, DM also shows two highly deviant experimental properties, absent in either L546A or L754A: these are an enormous primary deuterium room temperature isotope effect of $661 \pm 27$ and its very low sensitivity to temperature ($\Delta E_r \sim 0$).$^{28,30}$

**DM as an Outlier: Enormous Temperature-Independent KIEs Can Be Rationalized by Modifications in Both the Hydrogenic Donor–Acceptor Distance (DAD) and the Frequency of DAD Sampling.** Based on the nonadiabatic PCET model that fits the temperature independent KIEs in many native enzymes,$^{54,60}$ the observed rate constants of SLO are determined by a conformational sampling coordinate and PCET reaction coordinate, which can be visualized in Figure 5 and formalized as eq 1.$^{9,28,33}$

$$k_{\text{obs}} = k_{\text{eq}} \cdot k_{\text{PCET}}$$

where $k_{\text{eq}} < 1$ represents the small fraction of active enzyme–substrate protein substrates that arise from stochastic sampling processes (Figure 5A), multiplied by $k_{\text{PCET}}$, the nonadiabatic rate constant of the H-tunneling process (Figure 5B–D). The $k_{\text{PCET}}$ is derived from a formulation that resembles the Marcus analysis for electron tunneling (including reorganization energy, $\lambda$, and driving force $\Delta G^\circ$ parameters), in which the barrier-less hydrogenic wave function overlap coordinate depends on a temperature-dependent coordinate for achieving transient degenerate energy levels between reactant and

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Figure 5. Rate constants for enzyme catalyzed nonadiabatic hydrogen transfer can be formalized as $k_{\text{obs}} = k_{\text{eq}} \cdot k_{\text{PCET}}$. (A) $k_{\text{eq}}$ ($\propto 1$) represents a stochastic ground state search through inactive conformations (b) to reach catalytically active E-S complexes (a conformer). Subsequent thermal sampling further reduces the distance between the H-donor and acceptor as seen in the pretunneling $a^\circ$ conformer. (B-D) illustrate contributions from $k_{\text{PCET}}$. (B) Heavy atom protein motions produce transiently degenerate energies for the reactant and product wells, a prerequisite for wave function overlap at the tunneling ready state (TRS) (C). (D) The effective potential along the DAD sampling coordinate varies, starting with $\Delta E_r \approx 0$ for native enzyme (top), becoming $\Delta E_r \gg 0$ when the DAD is elongated following single mutations (middle) and finally arriving at the catastrophic DM scenario in which the DAD is elongated, but coupled with a rigid DAD sampling potential (bottom). In panel D, light purple refers to the shorter wave function distribution for deuterium, and dark purple represents the more distributive protium wave function.
product (Figure 5B,C). The addition of a thermally averaged DAD sampling mode completes the analysis (Figure 5D).

The latter has emerged as an important ruler for the degree of active site optimization and can be quantitated by the magnitude of $\Delta E_a$.

In naturally evolved (i.e., WT) enzymes, active sites for enzymatic C–H activation reactions are finely tuned by way of conformational sampling, to achieve highly compacted DADs in pretunneling configurations, labeled conformer (a*) (Figure 5A). When this occurs, the coordinate in Figure 5B produces the tunneling ready state (TRS) (Figure 5C) via thermal excitation that is largely independent of the labeling of substrate with deuterium; in this instance, further optimization of the reaction coordinate via DAD sampling (Figure 5D) is either absent or minor. The DAD sampling coordinate becomes increasingly important for impaired enzymes systems, leading to an incremental increase for $E_{a(D)}$ relative to $E_{a(3)}$, and is easily detected from an increase in $\Delta E_a$ away from a WT value that is generally small, close to zero. The isotope dependence of the DAD sampling coordinate, Figure 5D, is often perceived as puzzling, but can be readily understood in the context of the smaller, less diffuse wavelength for deuterium that introduces a requirement for shorter DADs; this can become an important source of the increased energetic barrier for achieving deuterium wave function overlap when the active site environment has been perturbed away from optimal alignment. The fact that mutation most generally also leads to softer active site potential energy surfaces enhances access to the DAD sampling process.

Historically, in the case of all catalytically impairing single mutants of SLO that include L546A and L754A, the $\Delta E_a$ increases relative to WT. However, the DM kinetic isotope effect is almost completely temperature independent ($\Delta E_a \sim 0$). While this would at first glance suggest a finely tuned active site, the significantly decreased $k_{cat}$ and enormous primary $Dk_{cat}$ of $\sim$700 of DM can only be ascribed to an elongated, nonoptimal DAD in the pretunneling conformer that prevents subsequent DAD sampling. When seen in this context, the collective kinetic data indicate that DM is simply unable to recover from a nearly catastrophically elongated DAD. Though the multiple biophysical probes in the present study were initially pursued with the goal of detecting and defining which aspects of protein structure/dynamics would be unique to DM and account for an implied rigidification, we have found no evidence for impairment to the dynamical network in L754A, which looks identical to WT in this regard. While a dynamical impairment does arise in the other component of DM, L546A, it alone is not serious enough to prevent DAD sampling and allows recovery to an active site configuration and KIE similar to WT. In the context of the behavior of these two parent mutants the most direct interpretation of the origin of the impairment in DM is seen as a combination of a serious mispositioning of substrate (due to L754A) that has simultaneously made it much more difficult to rearrange itself into a configuration that permits tunneling (due to the additional impairment of the protein dynamical network from L546A).

DM as an Outlier: Mispositioning of the Reactive Carbon of Substrate into a Region of SLO That Lies Outside of the Native Protein Dynamical Network. In light of the current findings, results from other methods were sought as a source of structural insight into the DM’s unique behavior. ENDOR studies have shown themselves to be a powerful probe for E-S complexes of SLO, enabling a direct interrogation of the dominant ground state H-donor and H-acceptor distances. Completed studies of WT show two such ground state structures, with one configuration (a) showing van der Waals distance for the hydrogen donor and acceptor and a second, inactive configuration (b) with a DAD almost 1 Å longer. Proficient wave function overlap in WT enzyme is proposed to arise via thermal sampling within the potential energy well of the “a” conformer to form DADs reduced from van der Waals (a*) (cf. Figure 5).

Importantly, ENDOR of the DM reveals only a single ground state “b” configuration that corresponds to the more extended, catalytically inactive form of E-S, consistent with the sluggish rate of this variant. In the same experiment, a single “b” configuration was also obtained with I553G, a mutation that like DM is predicted to exhibit an elongated DAD, but unlike DM, has a catalytic activity only 5-fold reduced from WT; the latter is accompanied by a highly temperature dependent KIE ($\Delta E_a \sim 5$ kcal/mol) that reflects active dynamical sampling back to a WT-like configuration, indicating that ENDOR alone cannot provide the elusive source of impairment for DM SLO. MD simulations were initiated for the ground state E-S structures of WT, DM and I553G using the metal to C11 and metal to C10 distances restrained by ENDOR measurements; significantly these studies suggested differences in the positioning of substrate within the DM form of SLO relative to WT and I553G.

Pursuing this possibility, a recent QM/MM study of the E-S complexes for WT and DM SLO provides a potential glimpse into the molecular origins for the collective impairment of mutations at both L546 and L754. QM/MM models have the advantage over the restrained MD simulations, because these calculations do not require any input from experimental findings to generate active site conformational. Corresponding ground state models of the active site structures for WT and DM derived from these computations show an almost superimposable C1–C7 region of LA that subsequently diverges in the region of substrate that encompasses the reactive carbon and its flanking double bonds (C9–C13). Importantly, the computed ground state DADs between C11 of LA substrate and metal-bound oxygen, of 3.2 Å for WT and 3.6 Å for DM, are remarkably consistent with the distances estimated from ENDOR experiments of 3.1 and 3.8 Å, respectively. It has been noted that the two altered side chains in DM, L546 and L754, are highly conserved and act to clamp the C11 of LA in position for reaction with the adjacent Fe(III)–OH. It is, therefore, of considerable interest that simultaneous replacement of both leucine side chains for alanine in SLO leads to a new QM/MM-based configuration for C11 of substrate, which has moved into the space normally filled by L754 (Figure 6).

A second observation in support of substrate mispositioning in DM comes from the accompanying QM/MM computation of $\Delta G^\circ$ for conversion of bound substrate to product. For DM, this $\Delta G^\circ$ is calculated to be more uphill by 10 kcal/mol relative to WT, in the same direction as the experimentally observed increase in the energy of activation of $\sim$10 kcal/mol for DM. Given an improbable ability of the SLO active site to alter the innate bond dissociation energy of bound substrate, together with our failure to detect significant changes in redox potential of the Fe cofactor in DM, a change in chemical driving force is ruled out as the source of the increased $\Delta G^\circ$ for DM. The most reasonable interpretation is a stabilization of substrate
within a binding mode that is highly compromised with regard to its distance and ability to promote wave function overlap from the C11 of substrate to the Fe(III)–OH acceptor. One caveat regarding the referenced QM/MM study is its inability to capture the experimental evidence for irreversible formation of pentadienyl radical under single turnover, anaerobic conditions.20,25 Scheme 2 suggests a way to rationalize the discrepancy between computation (ΔG° > 0) and experimental findings (ΔG° < 0). As shown, completion of the DM reaction will entail a return of the initially formed radical intermediate (illustrated in Scheme 1) to a less active but thermodynamically more stable binding position (E-I). This leads to the net exergonic process observed experimentally for the conversion of E-S to E-I. In this context, the trajectory captured by QM/MM for DM represents only a portion of the reaction in which E-S has been converted to E-I via an uphill process.

Figure 6. Comparison between the ground state substrate binding mode in WT (blue) and DM (yellow). The reactive carbon C11 is colored as black, iron colored as orange.


- **CONCLUSION**

The paper presents the biophysical characterization of a kinetically impaired double mutant of soybean lipooxygenase L546A/L754A and its parent single mutants L546A and L754A. Beyond adopting features from both parent single mutants, DM demonstrates a distinctive substrate binding mode that mispositions the substrate into configurations that are unfavorable for hydrogen tunneling. In the WT SLO, L546 resides within a protein motional network communicating between the surface loop and active site (Figure S2).23 It is of considerable interest that while L546A alone disrupts the protein motional network connecting the catalytically relevant surface loop and the active site,23 the enzyme is still able to undergo DAD sampling to achieve a KIE similar to WT.24 The movement of C—H of substrate into the “hole” created by the second mutation of L756 in DM has the additional effect of removing the reactive carbon of substrate from access to such thermal activation. This combination of a new substrate binding mode that is removed from access to thermally activated protein modes provides an explanation for the divergent features of DM that include its greatly impaired kcat, excessively elevated energy of activation and its enormous and close to temperature independent KIE. The current study, thus, provides a critical examination and insightful structural example of how relatively minor alterations in the placement of the reactive bond(s) of a bound substrate can cause dramatic alterations in the efficiency of enzymatic rate acceleration. In this context, the findings are highly relevant to the generic origins of substrate specificity in enzymes, whereby changes in the volume of active site side chains may either activate or impair an intended reaction course.

- **MATERIALS AND METHODS**

**Materials.** All reagents were purchased from commercial sources and used without further purification unless otherwise indicated. Wild-type and mutant SLOs were prepared, expressed and purified according to the previous report.26,29,31

**EPR Sample Preparation.** The EPR samples were prepared according to the previous protocol30 with several modifications. Each sample contained 140 μM SLO and 1 equiv of respective catechols in 0.1 M potassium phosphate (pH = 7.0). The ferric (yellow) lipooxygenase was first prepared from the mixing purified ferrous lipooxygenase and linoleic acid. The concentrated, purified ferrous lipooxygenase (300 μM) was titrated with linoleic acid (10 mM) at 4 °C, 0.1 M borate buffer (pH = 9.0) while monitoring the optical spectrum between 300 and 800 nm for development of the 330 nm peak characteristic of the ferric lipooxygenase. For WT and mutants, 2 equiv of linoleic acid was required to fully oxidize the protein. The ferric lipooxygenase (ca. 280 μM) was removed from the cuvette and dialyzed overnight in 0.1 M potassium phosphate buffer (pH = 7.0) at 4 °C. The sample was then stored at −80 °C until use. A known amount of catechols was transferred to an anaerobic glovebox, then dissolved in deoxygenated 0.1 M potassium phosphate with a final concentration of 280 μM. The catechols solution (100 μL) was mixed with deoxygenated ferric lipooxygenase (100 μL) and transferred into a suitable EPR tube. The tube was sealed with rubber stopper and kept at 4 °C. After 12 h, the tube was removed from the glovebox and frozen in liquid nitrogen within 1 min of its removal. The samples were stored at liquid nitrogen temperature until data acquisition. For each SLO variant, two control EPR samples were prepared as described above but with minor variations: One only contains 140 μM ferric lipooxygenase; another only contains 140 μM ferrous lipooxygenase.

**EPR Spectroscopy.** Samples for X-band (∼9.4 GHz) EPR spectroscopy were measured at the CalEPR center at the University of California, Davis. Continuous wave (CW) spectra were collected.
were removed from the immobilized pepsin with spin cups and the digestion in water) were previously determined for WT SLO.33 Samples were stored at atmospheric moisture condensation. The column exit was connected with 0.32 M citric acid stock solution at 0°C and acid quenched (to pH 2.4, combined with pH electrode). Samples were incubated randomly at mass resolution setting of 100 000 (at 350 to 1800 using the Orbitrap mass analyzer, in pro.

The values were normalized for 100% D2O and from this stock sample were diluted 10-fold in D2O. Each sample was thawed immediately prior to injection onto the column. The elution program consisted of a linear gradient from 5% B for 5.5 min, at a linear gradient to 5% B over 0.5 min, and isocratic conditions at 1.0 mm inner diameter, 5 μm particles, Restek, Bellefonte, PA) and guard precolumn (C8, Restek). Solvent A was 99.9% water/0.1% formic acid and solvent B was 99.9% acetonitrile/0.1% formic acid (v/v). Each sample was thawed immediately prior to injection onto the column. The elution program consisted of a linear gradient from 5% to 10% B over 1 min, a linear gradient to 40% B over 5 min, a linear gradient to 100% B over 4 min, isocratic conditions at 100% B for 3 min, a linear gradient to 5% B over 0.5 min, and isocratic conditions at 5% B for 5.5 min, at a flow rate of 300 μL/min. The column compartment was maintained at 4°C and lined with towels to absorb atmospheric moisture condensation. The column exit was connected to the ESI source of the mass spectrometer using PEEK tubing (0.005″ inner diameter × 1/16″ outer diameter, Agilent). Mass spectra were acquired in the positive ion mode over the range m/z = 350 to 1800 using the Orbitrap mass analyzer, in profile format, with a mass resolution setting of 100 000 (at m/z = 400). Data acquisition was controlled using Xcalibur software (version 2.0.7, Thermo).

HDX Data Analysis. Mass spectral data acquired for HDX measurements were analyzed using the software, HDX WorkBench. The percent deuterium incorporation was calculated for each of these peptides, taking into account the number of amide linkages (excluding proline residues) and the calculated number of deuterons incorporated. The values were normalized for 100% D2O and corrected for peptide-specific back-exchange, as determined previously.31 The data were plotted as deuterium exchange versus time using Igor Pro software. The rates and extents of exchange (at 4 h) were determined from one- or two-exponential fits to the analyzed time-resolved HDX data. The weighted average rates of HDX were determined from the individual rate constants of these two exponentials. Apparent enthalpies of activation from HDXMS rates (Ea[HDX(avg)]) were determined for each peptide and mutation from the linear fits to the Arrhenius plots of the HDXMS weighted average exchange rates versus inverse temperature. These Ea[HDX(avg)] values represent the mean ± s.d. from the five temperatures studied here.

Room-Temperature Crystallography. All additional protein purification and crystallization steps were performed as previously described.33 Diffraction data were collected at room temperature at Beamline 8.3.1 of the Advanced Light Source at Lawrence Berkeley National Laboratory. Data were processed by the ELVES program,28 with integration performed in MOSFLM26 and scaling and merging in SCALA, POINTLESS and TRUNCATE,67 or using XDS2 with scaling and merging using AIMLESS,27 POINTLESS and TRUNCATE in the CCP4 suite provided through SGBGRID.77 An initial solution was found by molecular replacement using Phaser with PDB 3PZW as the search model. Manual refinement was performed in Coot and automated refinement using the PHENIX suite. Alternate conformers were modeled using qFit.76 We manually deleted waters that did not fit into the Fo-Fc map density and edited alternate side chain conformations as indicated in the positive density of the Fo-Fc map. Active site volumes were calculated using CASTp. The relative size of the pockets in each mutant was based on analysis of the most populated conformer of the residues lining the pocket and included residues Glu495, Leu496 His499, Trp500, His504, Ile538, Ala542, Leu546, Ile547, Ile553, Phe557, Ser567, Ser567, Val570, Ile571, Leu574 and Ile589. See Table S4 for data processing and refinement statistics.

ASSOCIATED CONTENT

Supporting Information

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Notes

The authors declare no competing financial interest.

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