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Local Electric Fields as a Natural Switch of Heme-Iron Protein Reactivity

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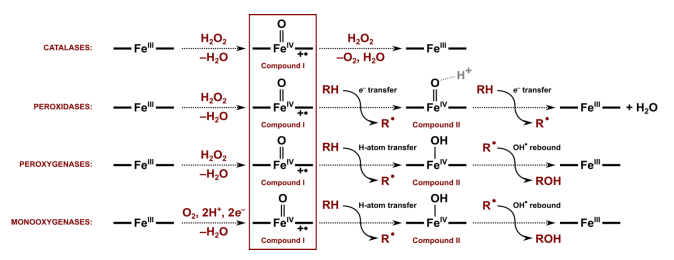
ABSTRACT

Heme-iron oxidoreductases operating through the high-valent Fe^{IV}O intermediates perform crucial and complicated transformations, such as oxidations of unreactive saturated hydrocarbons. These enzymes share the same Fe coordination, only differing by the axial ligation, e.g., Cys in P450 oxygenases, Tyr in catalases, and His in peroxidases. By examining ~200 heme-iron proteins, we show that the protein hosts exert highly specific intramolecular electric fields on the active sites, and there is a strong correlation between the direction and magnitude of this field and the protein function. In all heme proteins, the field is preferentially aligned with the Fe–O bond (F_z). The Cys-ligated P450 oxygenases have the highest average F_z of 28.5 MV cm⁻¹, i.e., most enhancing the oxyl-radical character of the oxo group, and consistent with the ability of these proteins to activate strong C-H bonds. In contrast, in Tyr-ligated proteins, the average F_z is only 3.0 MV cm⁻¹, apparently suppressing single-electron off-pathway oxidations, and in His-ligated proteins, F_z is -8.7 MV cm⁻¹. The operational field range is given by the trade-off between the low reactivity of the Fe^{IV}O Compound I at the more negative F_z , and the low selectivity at the more positive F_z . Consequently, a heme-iron site placed in the field characteristic of another heme-iron protein class loses its canonical function, and gains an adverse one. Thus, electric fields produced by the protein scaffolds, together with the nature of the axial ligand, control all heme-iron chemistry.

Keywords: Electric Field, Enzymes, Ferryl, Heme Iron, Redox Chemistry

INTRODUCTION

High-valent metal-oxo and metal-hydroxo intermediates are irreplaceable in Nature, as they can catalyze energetically demanding transformations, such as oxidations of (inactivated) saturated hydrocarbons.^{1–4} Heme-iron proteins may serve as an example using molecular oxygen or hydrogen peroxide to form the reactive species, known as Compound I (CpdI), to oxidize or oxygenate diverse organic substrates (**Scheme 1**).^{5–8} Particularly, hydroxylation of the substrates triggered by the cytochrome P450 family of proteins represents a cornerstone reactivity in xenobiotics' metabolic pathways, making them more soluble and more susceptible for elimination. Although debate remains on which factors dominate (or significantly contribute to) the reactivity of these systems,^{9–13} the activation barrier is indisputably governed by their ability to activate the strong C–H bonds of the substrates. C–H bond-dissociation free energies (BDFEs) are ca. 80-110 kcal mol⁻¹ and p K_A s are above 50,^{14,15} making the observed rates for transformations all the more impressive, in the range of $10^7 M^{-1} s^{-1.16}$



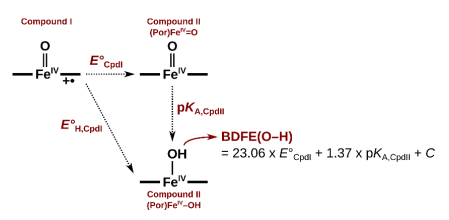
Scheme 1. Typical reactions performed by different classes of the heme-iron oxidoreductases. In the absence of a substrate, the resting state of the enzymes is the ferric (Fe^{III}) with a bound water molecule. The resting state is activated either by *(i)* Fe^{III} binding hydrogen peroxide, leading ultimately to the (Por⁺⁺)Fe^{IV}=O CpdI intermediate with the release of an H₂O molecule, or *(ii)* Fe^{III} reduction, followed by the Fe^{II} binding of molecular oxygen, leading to the same (Por⁺⁺)Fe^{IV}=O CpdI intermediate through sequential one-electron two-proton transfer and H₂O molecule release. Although the nature of the CpdI intermediate is the same, the electronic structure and reactivity may differ as a result of the nonidentical protein environment, such as axial ligands (Cys, His, or Tyr) and the protein fold. In the catalases enzymes, the CpdI reacts with the second molecule of H₂O₂ to yield O₂ and H₂O. In the peroxidases and oxygenases subclasses, the second reactive intermediate – the (Por)Fe^{IV}=O CpdII or

(Por)Fe^{IV}–OH CpdII – is proposed to be formed *via* single-electron transfer (associated with coupled proton transfer from substrate to some nearby amino acid residue) or electron-proton transfer to the CpdI. Lastly, CpdII can also oxidize or oxygenate the organic substrate.

Electric fields have been proposed to play a role in enzymatic catalysis. For example, Warshel went as far as postulating that the catalysis by enzymes is in general empowered by electrostatic preorganization: the placement of charged groups throughout the protein in a way that permanently electrostatically favors the transition state and disfavors the reactants, thus lowering the reaction barrier via a mechanism inaccessible in solution.^{17–19} Electrostatic preorganization or, more specifically, the role of electric fields in enzymatic (and non-enzymatic) catalysis has been vigorously investigated, probed, and debated in recent years, and in enzymology the focus thus far has been on a limited number of systems, such as ketosteroid isomerase and artificial Kemp eliminases.^{20–29} Given the key role of the Fe^{IV}-oxo bond in CpdI and CpdII in the heme-iron catalysis, the well-defined, rigid structure of the heme site presents but a perfect opportunity to apply an intermolecular field to maximize or control the reactivity. The electric field applied to Fe^{IV}O may, for example, give the Fe^{IV}-oxo bond a stronger oxyl-radical (Fe^{III}O⁻⁻) character, which is often associated with the higher metal-oxo electrophilicity.^{12,30–34}

Indeed, the oriented external electric field (OEEF) parallel to the Fe–O axis was proposed to induce changes in reactivity and selectivity of several heme iron-oxo CpdI models outside of the protein.^{35–40} Such an OEEF can significantly perturb the electronic structure of CpdI: localization of the unpaired electron in its ground state, and the spin-state (doublet/quartet) ordering and energy splitting, impacting the physicochemical properties.³⁹ In particular, the sign and the magnitude of the OEEF along the Fe–O axis were evidenced to alter the location of the radical site: on the nonbonding a_{1u}/a_{2u} orbital on porphyrin (denoted as Por⁺⁺), or on the adjacent Trp residue (denoted as Trp⁺⁺), in a pair of His-ligated peroxidases – ascorbate peroxidase (APX) and cytochrome *c* peroxidase (C*c*P).³⁹ Note that in Nature, these two peroxidases have different electronic structures, and the ultimate factor differentiating between the Por⁺⁺ and Trp⁺⁺ radicals has not been plausibly clarified.⁶ However, QM/MM calculations of CpdI in *CcP* and APX correctly reproduced the differences in the electronic structures of their identical QM regions, apparently affected by the protein edifice.^{41,42} From these findings, the intramolecular local electric fields (LEF) transpire as a strong and thus far not rigorously questioned suspect governing the reactivity and selectivity in the natural heme-iron proteins.

Still using a small model system representing the Cys-ligated CpdI of cytochrome P450, Shaik et al. proposed that the selectivity between the epoxidation and hydroxylation competing pathways could be switched by altering the direction and magnitude of the OEEF.⁴⁰ A sufficiently large OEEF of ca. $F = \pm 0.0125$ a.u. (~ ± 64 MV cm⁻¹) orthogonal to the porphine plane were calculated to realize a 100 % selectivity toward C–H vs. C=C bond activation.⁴⁰ As of now, it is unclear whether such dramatic fields can be achieved inside the heme-iron proteins. However, it is clear that, since OEEF affects the oxyl-radical character, it should impact the thermodynamic properties of the CpdI and CpdII, such as BDFE(O–H) of CpdII, their electron affinities (reduction potentials), and proton affinities (pK_As) (**Scheme 2**). Although the overall picture of the high-valent metal-oxo reactivity is extraordinarily convoluted, it can be strongly hypothesized that LEF, on-par with the nature of the axial ligand and its hydrogen-bonding pattern, likely substantially contributes to the oxidative powers and selectivity of CpdI and CpdII, and hence the rates of individual transformations.



Scheme 2. Interconversion of the oxygenated heme-iron CpdI and CpdII intermediates, and the associated thermodynamic quantities. Note that BDFE(O–H), equivalent to $E^{\circ}_{H,CpdI}$, quantifies an H-atom-abstraction driving force mediated by the CpdI Fe^{IV}O intermediate. Considering the thermodynamic equivalence of the two pathways connecting Fe^{IV}O CpdI and Fe^{IV}-OH CpdII, the BDFE(O–H) can be expressed by E°_{CpdI} and $pK_{A,CpdII}$; C is a standard reduction potential of H⁺/H⁺ in a particular solvent (C=57.6 kcal mol⁻¹ in water).

In this work, we first aim to elucidate what kinds of LEF (directions and magnitudes) the hemeiron sites experience inside the various natural oxidoreductases. We find a wide variety of LEFs, but such that they appear to correlate with the functions of these proteins, and specifically the reactivity of the presumptive CpdI or CpdII intermediates. Next, using the Cys-ligated cytochrome P450 monooxygenases, Tyr-ligated catalases, and His-ligated peroxidases as prototypical examples, we demonstrate that LEF steers the selectivity toward the productive substrate oxidation. Our findings infer that LEFs are indeed evolutionarily altered in heme-iron proteins to stir their reactivity, thus contributing to the understanding of the fundamental aspects of proteins operating through the metaloxo intermediates.

RESULTS AND DISCUSSION

The local electric fields in the active sites of heme-iron oxidoreductases

We analyzed the crystal structures of ca. 200 individual Fe-containing heme-based oxidoreductases (EC number: 1): catalases, cytochrome P450 monooxygenases and peroxygenases, heme oxygenases, peroxidases, and others, and elucidated the direction and magnitude of the LEF at their Fe centers (see **Computational Details** section in the SI). Note that the evaluated fields are induced by the entire protein *outside* of the heme-iron site, i.e., excluding the effects of the site itself and the axial ligand. The axial ligands thus *do not* contribute to the discussed LEFs. Instead, the LEFs, as presented in the article, precisely correspond to the OEEFs previously applied to the whole 'axial ligand–porphine–FeO' models, and thus affect the reactivity on top the effect of the axial ligand. The electric fields in the *x*,*y*,*z* directions (*cf.* **Figure 1**) are summarized for each protein in **Table S2** in the SI, whereas the proteins, grouped according to their EC numbers and the nature of their axial ligands, are listed in **Figures 2-4** and **Table 1**.

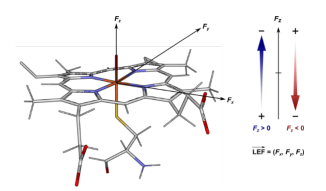


Figure 1. The orientation of the components of the electric field vector in the heme-iron model. The F_x and F_y components are oriented in the porphine plane and the Fe–N_A and Fe–N_B bonds of the porphyrin. The F_z component is orthogonal to the porphine plane, parallel with the Fe–O bond of the putative CpdI species.

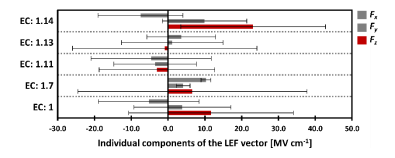


Figure 2. The average values of the F_x , F_y , and F_z components of the LEF vector among the subclasses of the oxidoreductase proteins (EC: 1). For each class, the LEF is displayed from top to bottom as the average F_x , F_y ,

and F_z components of the LEF vector in MV cm⁻¹ shown as thick bars (F_z is highlighted in red), and the thin black lines represent the standard deviations, σ .

A marked preference for the LEF oriented in the z direction across the whole family of proteins is evident from Figure 2. The average values of LEF are $F_x = -5.2$ MV cm⁻¹, $F_y = 3.9$ MV cm⁻¹, $F_z =$ 11.7 MV cm⁻¹. A higher preference for the z orientation of the LEF is found for the proteins in the EC subclasses 1.7 (oxidoreductases acting on other nitrogenous compounds as donors), 1.13 (oxidoreductases acting on single donors with incorporation of molecular oxygen (oxygenases)), and 1.14 (oxidoreductases acting on paired donors, with incorporation or reduction of molecular oxygen), indicating a more distinct regulation of the enzymes' catalytic power by the LEF, as compared to an electrostatically featureless, homogeneous environment. Note that proteins in the EC classes 1.7 and 1.13 have small average F_z (Figure 2), but this is due to the equally large preference for the high positive and high negative F_z (as embodied in the substantial standard deviations of the F_z in Figure 2). The highest average F_z is thus found in the subclass of EC 1.14: $F_x = -7.5$ MV cm⁻¹, $F_y = 9.9$ MV cm⁻¹, and $F_z = 23.1$ MV cm⁻¹. This should come as no surprise, as this class comprises mostly the cytochrome P450 oxygenases, which activate the strong C-H bonds of organic substrates. The mostly positive F_z in this class of proteins (*cf.* Figure 3) adequately indicates a raised oxygenation reactivity, and thus strongly suggests the careful tuning of the LEF by the evolution. A large standard deviation of the F_z (in the EC: 1.14 class, the $\sigma(F_z) = 19.7$ MV cm⁻¹) further suggests a fine-tuning of the proteins' reactivity or selectivity of the individual proteins through F_z . As a prominent example, the cholesterol 7-alpha-hydroxylase (CYP7A1; PDB code: 3v8d), an important enzyme in the metabolism of cholesterol putatively operating through the CpdI intermediate, attains a strong F_z of 55.8 MV cm⁻¹ (in contrast with the relatively weak $F_x = -19.5$ MV cm⁻¹, and $F_y = 14.6$ MV cm⁻¹; cf. Table 1).

On the contrary, the oxidoreductases that act on the peroxide (EC 1.11; *cf.* Figure 2) show a much lower magnitude of the LEF vector with less strictly induced orientation parallel with the Fe–O axis. The average LEF components are: $F_x = -4.6$ MV cm⁻¹, $F_y = -3.5$ MV cm⁻¹, and $F_z = -3.0$ MV cm⁻¹. Besides, F_z is preferably negative in the individual EC: 1.11 subclasses (Figure 3), and that is consistent with a lower reactivity of their CpdI intermediates that *are not* involved in the H-atom abstractions. In other words, lower F_z is related to a lower CpdI reactivity. This can be supported by the fact that the CpdI intermediates in catalases and peroxidases (both in EC: 1.11.1) were found to be more stable, and as such, more readily isolated in the past than CpdI intermediates in the P450 proteins.⁴³

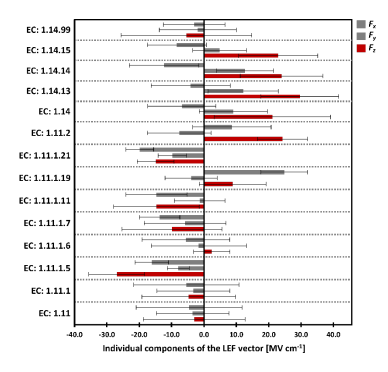


Figure 3. The average values of the F_x , F_y , and F_z components of the LEF vector among the subclasses of the oxidoreductase proteins: *(i)* acting on a peroxide as acceptor (EC: 1.11) and *(ii)* acting on paired donors, with incorporation or reduction of molecular oxygen (EC: 1.14); average values are shown as thick bars, and thin lines represent standard deviations, σ .

Importantly, in all of the proteins examined herein, there are only three different axial ligands present – Cys, Tyr, and His, and with certain simplification, we may associate each of the ligands with a specific protein function from **Scheme 1**. The catalase reactivity (EC: 1.11.1.6) is mostly associated with the Tyr-axial heme-iron ligation, the peroxidase reactivity (EC: 1.11.1., except for 1.11.1.6) - with the His ligation, and the oxygenase/peroxygenase reactivity (EC: 1.11.2. and 1.14, except for 1.14.99) - with the Cys ligation. **Figure 4** reveals a pronounced correlation between the proteins LEFs and their function, explicitly evaluating the heme-iron axial ligation (see also **Table 1**). In particular, the preference for the *z* component of the LEF is increased in the Tyr – His < Cys series (**Figure 4**); while the Tyr-ligated proteins have a preference for slightly positive F_z , His-ligated proteins exhibit slightly negative F_z , and Cys-ligated proteins exhibit strictly positive and large F_z (**Figure 4**).

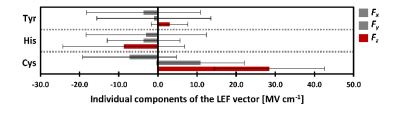


Figure 4. The average values of the F_x , F_y , and F_z components of the LEF vector among the oxidoreductase proteins (EC: 1) with Tyr-, His-, and Cys-axial ligation; average values are shown as thick bars, and thin lines represent standard deviations, σ . Recall that the axial ligand itself is not included in the calculations of LEF; hence, the axial ligand adds to the effect depicted here.

In detail, the Cys-ligated proteins – consisting mostly of the P450 oxygenases (EC: 1.14) and peroxygenases (EC: 1.11.2) – on average exhibit the $F_z = 28.5$ MV cm⁻¹. From refs. ^{40,44}, such LEF can be expected to decrease the H-atom-transfer activation barrier by ca. 4 kcal mol⁻¹ owing to an increase in the oxyl-radical character of the Fe^{IV}O.

The Tyr-axial ligated proteins (i.e., catalases; EC 1.11.1.6), on the contrary, exhibit a very small field in the positive *z* orientation: $F_z = 3.0$ MV cm⁻¹. However, the field in the Tyr-ligated proteins, though minimal in the *z* direction, is not randomly oriented, either. Instead, the low F_z appears to be genuinely induced by the protein fold, as manifested by the small standard deviation of the $\sigma(F_z) = 4.7$ MV cm⁻¹, and compared with the standard deviation of the F_x and F_y components for the whole protein series: $\sigma(F_x) = 13.7$ MV cm⁻¹ and $\sigma(F_x) = 13.2$ MV cm⁻¹, which suggest a greater randomness.

| | Average F_x | | Average F_y | | Average F_z | |
|----------------------|------------------------|------------|------------------------|------------------------------|------------------------------|------------------------------|
| | $[MV \text{ cm}^{-1}]$ | | $[MV \text{ cm}^{-1}]$ | | [MV cm ⁻¹] | |
| All proteins | -5.2 ± 13.7 | | 3.9 ± 13.2 | | 11.7 ± 22.4 | |
| Cys-ligated | -7.2 ± 12.0 | | 10.9 ± 11.1 | | 28.5 ± 14.1 | |
| Examples: | PDB code | EC number | Function | F_x [MV cm ⁻¹] | F_y [MV cm ⁻¹] | F_z [MV cm ⁻¹] |
| P. putida P450cam | $1dz9^a$ | 1.14.15.1 | monooxygenase | -5.1 | -0.3 | 25.2 |
| S. coelicolor CYP158 | 2d09 | 1.14.19.69 | biflaviolin synthase | -9.4 | <i>8.3</i> | 25.7 |
| H. sapiens CYP7A1 | 3v8d | 1.14.14.23 | monooxygenase | -19.5 | 14.6 | 55.8 |
| Tyr-ligated | -ligated -3.7 ± 14.5 | | -1.0 ± 14.6 | | 3.0 ± 4.7 | |
| Examples: | PDB code | EC number | Function | F_x [MV cm ⁻¹] | F_y [MV cm ⁻¹] | F_z [MV cm ⁻¹] |
| M. luteus KatA | lgwf ^a | 1.11.1.6 | catalase | -11.0 | -17.9 | 4.1 |
| P. mirabilis KatA | 1 mqf ^a | 1.11.1.6 | catalase | -11.1 | -9.1 | 8.6 |
| H. pylori KatA | $2a9e^{a}$ | 1.11.1.6 | catalase | -21.2 | -10.5 | 1.7 |
| N. crassa KatA | 6nsw ^a | 1.11.1.6 | catalase | 12.2 | 19.7 | -1.3 |
| His-ligated | -3.0 ± 15.4 | | -3.7 ± 9.3 | | -8.7 ± 15.6 | |

 Table 1. The average components of the LEF vector in the oxidoreductase proteins grouped according to the nature of the heme-iron axial ligand. The individual components of the LEFs are indicated for several examples in each of the groups, as discussed in the text.

| Examples: | PDB code | EC number | Function | F_x [MV cm ⁻¹] | F_y [MV cm ⁻¹] | F_z [MV cm ⁻¹] |
|----------------------|--------------------------|-----------|-------------------------|------------------------------|------------------------------|------------------------------|
| S. cerevisiae CcP | <i>lebe</i> ^a | 1.11.1.5 | peroxidase | -22.1 | -4.8 | -26.7 |
| A. rusticana HRP | 1hch ^a | 1.11.1.7 | peroxidase | -5.5 | -9.4 | -36.6 |
| G. max APX | 5jqr ^a | 1.11.1.11 | peroxidase | -4.4 | 9.0 | 4.1 |
| H. sapiens MPO | 5fiw | 1.11.2.2 | peroxygenase | -9.1 | -3.1 | 28.4 |
| C. hircus LPO | 3rke | 1.11.1.7 | peroxidase | -15.1 | -23.1 | 12.4 |
| A. ramosus ARP | 2e39 | 1.11.1.7 | peroxidase | -3.9 | -1.1 | -2.1 |
| L. major LmP | 3riv | 1.11.1.11 | peroxidase | -11.5 | -9.7 | -15.5 |
| B. pseudomallei KatG | $5sx0^a$ | 1.11.1.21 | catalase- peroxidase | -17.5 | -9.2 | -15.0 |

^aProteins were crystallized in the CpdI or CpdII form.

Finally, the His-ligated proteins contain mostly heme oxygenases (EC: 1.14.99) with the average LEF components of $F_x = -3.3$ MV cm⁻¹, $F_y = -2.1$ MV cm⁻¹, $F_z = -5.9$ MV cm⁻¹, dioxygenases (EC: 1.13.11) with $F_x = 5.2$ MV cm⁻¹, $F_y = -2.2$ MV cm⁻¹, $F_z = -9.5$ MV cm⁻¹, and peroxidases (EC: 1.11.1; except for 1.11.1.6) with $F_x = -8.0$ MV cm⁻¹, $F_y = -5.9$ MV cm⁻¹, $F_z = -9.5$ MV cm⁻¹. Noticeably, there is a certain preference for the negative F_z (ca. 71 % of a total of 70 studied Hisligated proteins exhibit negative F_z ; see Table S2 in the SI), contradicting the higher oxyl-radical character of the CpdI. Recall that F_z in the Cys-ligated proteins (such as the P450 oxygenases/peroxygenases) is positive in 100 % of 95 of the studied cases. F_z in the His-ligated proteins ranges from $F_z = -37$ MV cm⁻¹ to $F_z = +49$ MV cm⁻¹, thus, suggesting a possibility of tuning the CpdI reduction potentials both in a positive and negative direction over a large total range of ca. 500 mV (vide infra), which might be important for activation (oxidation) of a manifold of substrates. This apparent tuning of redox properties is illustrated in Figure 5 (left), correlating the experimental E°_{cpdI} values of various His-ligated peroxidases⁴⁵ and their LEF F_z established in this work. Specifically, myeloperoxidase (MPO) has $E^{\circ}_{cpdI} = 1.35$ V, and $F_z = +28.4$ MV cm⁻¹; ascorbate peroxidase (APX) has $E^{\circ}_{cpdI} = 1.16$ V, and $F_z = +4.1$ MV cm⁻¹; lactoperoxidase (LPO) has $E^{\circ}_{cpdI} = 1.14$ V, and $F_z = +12.4$ MV cm⁻¹; arthromyces ramosus peroxidase (ARP) has $E^{\circ}_{cpdI} = 0.92$ V, and $F_z = -2.1$ MV cm⁻¹; horseradish peroxidase (HRP) has $E^{\circ}_{cpdI} = 0.90$ V, and $F_z = -36.6$ MV cm⁻¹).

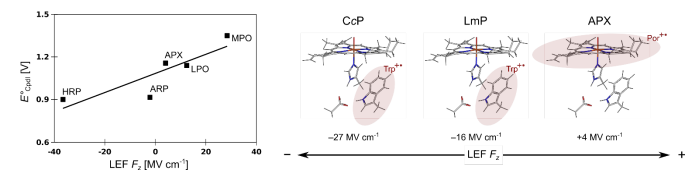


Figure 5. *Left:* Experimental E°_{cpdI} electrochemical potentials of various peroxidases plotted against the calculated F_z components of their LEFs. *Right:* Different electronic structure of the $(Trp^{++})Fe^{IV}O$ CpdI in CcP and LmP vs. $(Por^{++})Fe^{IV}O$ CpdI in APX as observed in Nature can be associated with the change in the proteins F_z component of the LEF vector. The unpaired radical is shifted from the Trp^{++} to Por^{++} as F_z becomes more positive. Note that we are not aware of equivalent experimental data for Cys- and Tyr-ligated oxidoreductases to be included for comparison.

At last, the cytochrome *c* peroxidase (C*c*P) and *leishmania major* peroxidase (LmP) exhibit LEF quite different from that in the ascorbate peroxidase (APX), which is otherwise nearly homologous at the active site (**Table 1**).[§] Since the two groups of enzymes have different electronic structures (e.g., the unpaired electron located on Trp⁺⁺ in C*c*P and LmP, and on Por⁺⁺ in APX; *cf.* Figure 5, *right*), this finding suggests LEF playing a role in tipping the balance between the two resonant forms. Noticeably, the higher F_z component of the LEF observed in the APX enzyme is indeed consistent with the relocation of the unpaired radical toward the Por⁺⁺ (*vide infra*).

The effect of LEF on the physicochemical properties of the CpdI and CpdII in the cytochrome P450 oxygenase, catalase, and peroxidase model systems

The ground spin state of the CpdI can be either quartet (S = 3/2) or doublet (S = 1/2), originating from the ferromagnetic or antiferromagnetic coupling of the 'Fe–O' triplet spin state and the unpaired electron located on (*i*) the porphine ring (Por), (*ii*) the axial ligand (Ax), or (*iii*) the adjacent Trp residue in case of the peroxidase model; *cf.* Figure 5, *right*. Employing the model systems constructed from the P450 monooxygenase (P450cam), catalase (from *P. mirabilis*), and peroxidase (APX) protein crystal structures (Figure S6), we next compare the electronic structure properties of their CpdI intermediates. In all of the models, density functional theory, DFT (B3LYP method; see Computational Details in the SI) predicts the quartet and doublet spin states to be nearly degenerate,

[§] The situation does not change by introducing the K⁺ cation in the APX enzyme, which is present in the crystal structure, but was omitted in the LEF analysis. We see almost no difference in the LEF component in the *z* direction (*cf.* $F_z = 4$ MV cm⁻¹ without the K⁺ ion vs. $F_z = 2$ MV cm⁻¹ with the K⁺ ion).

lying within ca. 1 kcal mol⁻¹ in free energy. In detail, the preferred configuration is calculated to be quartet in Cys- and Tyr-ligated models (by 0.7 kcal mol⁻¹ in both) and doublet in His-ligated model (by 0.3 kcal mol⁻¹), see also **Supporting Note S3** in the SI. It is evident from the Mulliken spin densities in **Table 2** that the radical in CpdI is preferentially located on the axial ligands (or Trp). However, it is also partially displaced to the porphine ring, and that is consistent with an operable mixing of the two resonance forms of the CpdI state, suggested to be a part of tuning of the CpdI properties in the P450 catalysis.^{46,47} Note that Mulliken spin densities can be only qualitative, due to the DFT delocalization error, which is known to overstabilize the radical-delocalized states.⁴⁸

A one-electron reduction of the $(Trp^{+*}/Ax^{+*}/Por^{+*})Fe^{IV}=O$ CpdI results in the formation of the Fe^{IV}=O CpdII in the triplet (*S* = 1) spin state in all of the cases, where the additional electron is paired with the Trp^{+*}/Ax^{+*}/Por^{+*} radical, not changing the electronic distribution between the Fe and O atoms significantly. In turn, this may be responsible for a small change in the Fe–O bond distance, as also observed in the experiments.⁴⁹ On the contrary, a transfer of an H atom (i.e., proton and electron pair) to CpdI yields a protonated Fe^{IV}-OH CpdII species with a significantly elongated Fe–O bond and electron density that is displaced from the O atom toward the Fe center (*cf.* **Table 2**).

| · / | 2 | ``` | | <i>,</i> |
|----------------------|--|--|--|--|
| | CpdI Fe ^{IV} =O (S = 1/2) | CpdI Fe ^{IV} =O (S = 3/2) | CpdII (Por)Fe ^{IV} =O (S = 1) | Cpd II (Por)F e^{IV} -OH (S = 1) |
| Cys-ligated P450 m | onooxygenase | | | |
| Fe [<i>e</i>] | 1.24 | 1.22 | 1.32 | 1.82 |
| O [<i>e</i>] | 0.82 | 0.85 | 0.81 | 0.16 |
| Por [<i>e</i>] | -0.33 | 0.24 | -0.11 | -0.14 |
| Cys [<i>e</i>] | -0.68 | 0.67 | -0.02 | 0.15 |
| Fe–O dist. [Å] | 1.65 | 1.66 | 1.69 | 1.84 |
| Tyr-ligated catalase | | | | |
| Fe [<i>e</i>] | 1.27 | 1.26 | 1.32 | 1.83 |
| O [<i>e</i>] | 0.78 | 0.82 | 0.77 | 0.13 |
| Por [<i>e</i>] | -0.33 | 0.18 | -0.10 | -0.13 |
| Tyr [<i>e</i>] | -0.72 | 0.74 | 0.01 | 0.17 |
| Fe–O dist. [Å] | 1.63 | 1.65 | 1.67 | 1.82 |
| His-ligated peroxide | ase | | | |
| Fe [<i>e</i>] | 1.26 | 1.25 | 1.28 | 1.90 |
| | | | | |

Table 2. The Mulliken spin densities of selected atoms and Fe–O bond distance of the CpdI and CpdII intermediates of the heme-iron P450 monooxygenase, catalase, and peroxidase models calculated at the DFT (B3LYP) level of theory in the zero field (see the **Computational Details** section in the SI).

| O [<i>e</i>] | 0.84 | 0.84 | 0.82 | 0.13 |
|------------------|-------|------|-------|-------|
| Por [<i>e</i>] | -0.41 | 0.27 | -0.09 | -0.05 |
| Trp [<i>e</i>] | -0.67 | 0.65 | 0.00 | 0.03 |
| Fe–O dist. [Å] | 1.65 | 1.66 | 1.66 | 1.81 |

The calculated $E^{\circ}_{H,CpdI}$ potentials of 1.28 V (Cys-ligated heme) and 0.88 V (His-ligated heme) are in excellent agreement with the measured potentials in refs. ^{45,50} (*cf.* Cys: $E^{\circ}_{H,CpdI} = 1.28$ V vs. 1.22 V for CYP158 in ref. ⁵⁰; His: $E^{\circ}_{H,CpdI} = 0.88$ V vs. ~0.9-1.2 V for various peroxidases with the same His-Asp-Trp/Phe triad in ref. ⁴⁵). Also, the calculated $E^{\circ}_{H,CpdI} = 1.18$ V of the catalase model seems to be realistic, as the electrochemical properties of the negatively charged Cys and Tyr axial ligands in the P450 and catalase enzymes were suspected to be of similar magnitude;^{51,52} however, since the CpdII is not a native intermediate in the catalase's enzymatic cycle,⁵³ the $E^{\circ}_{H,CpdI}$ has no practical (biological) relevance. The calculated values of the $E^{\circ}_{H,CpdI}$ potentials are also consistent with the highest reactivity of the CpdI in the P450 enzyme and correspond to a decreasing BDFE(O–H) of ca. 97 kcal mol⁻¹ (Cys) > 94 kcal mol⁻¹ (Tyr) > 88 kcal mol⁻¹ (His). We also note that the calculated proton affinities of CpdII are consistent with the trend in the experimentally obtained $pK_{A,CpdII}$; *cf.* the proton affinities of -316 kcal mol⁻¹ (Cys), -310 kcal mol⁻¹ (Tyr), and -293 kcal mol⁻¹ (His) vs. the $pK_{A,CpdII}$ of ~12 (CYP158)⁵¹, ~13 (*H. pylori* catalase)⁵², and ≤ 4.5 (APX)⁵⁴. Similarly, the electron affinities of CpdI give the opposite trend of -88 kcal mol⁻¹ (Cys), -91 kcal mol⁻¹ (Tyr), and -101 kcal mol⁻¹ (His).

By applying the oriented external electric field (OEEF) in the Fe–O direction, we next evaluate how the specific values of F_z actually attainable in the proteins impact the electronic structure and physicochemical properties of each proteins' CpdI. Since the individual protein classes differ significantly in function (*cf.* Scheme 1), only the heme-iron oxygenases that abstract H atoms from the substrates would have distinct transition states that we could calculate. We thus rather focus on the properties of the oxygenated intermediates, as they are unifying for all of the studied proteins. Also, the Fe^{IV}O CpdI are extremely reactive species and operate through the transition states that are very 'reactant-like'. The effect of the field in the Fe^{IV}O intermediate and the transition state thus should be comparable and can be related with heme-iron reactivity.

First, the OEEF in the positive Fe–O direction results in the relocation of the radical toward Por^{+•} (Figure 6), as was already described in ref. ⁴⁰. Besides others, this allows to quantify the effect of the LEF on the peroxidases proteins, C*c*P, LmP, and APX, and their electronic structure; e.g., the location of the radical in the CpdI intermediates. These peroxidases differ in LEF F_z by ca. 31 MV cm⁻¹ (going from $F_z = -26.7$ MV cm⁻¹ in C*c*P to $F_z = +4.1$ MV cm⁻¹ in APX). Correspondingly, from

applying the OEEF of the same magnitudes and directions, we predict the spin densities to be ca. $0.96/0.09 \ e$ on Trp/Por in both C*c*P and LmP, and $0.47/0.59 \ e$ on Trp/Por in APX. Therefore, ca. $0.5 \ e$ can be expected to transfer from Trp to Por due to the effect of LEF in these proteins. Note that higher radical delocalization might be observed as a result of the DFT delocalization error.⁴⁸ Also, the calculations are performed on the small model systems; however, effects beyond electrostatics that arise from the second-coordination shell may also contribute to the experimental observations.

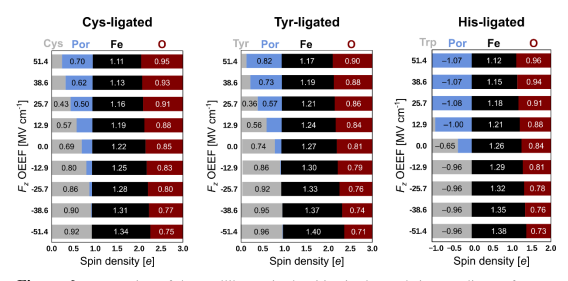


Figure 6. Progression of the Mulliken spin densities in the CpdI intermediates of Cys-, Tyr-, and His-ligated heme-iron models with the change of the F_z component of OEEF.

Additionally, the Fe–O oxyl-radical character is increased as the effect of a higher F_z (*cf.* the Mulliken spin density progression at the O atom in **Figure 6**), leading to a higher CpdI electrophilicity. In detail, we observed an increase in the $E^{\circ}_{H,CpdI}$ of several hundreds of millivolts when the F_z of +51.4 MV cm⁻¹ was applied (= 0.01 a.u.; roughly corresponding to the highest LEF observed in the heme-iron oxidoreductases enzymes; *vide supra*). Since the increase in the $E^{\circ}_{H,CpdI}$ is equivalent to an increase in the BDFE(O–H), this finding has an important implication for the reactivity of the heme-iron proteins, as it provides a recipe for how the proteins can utilize their sequences and folds to tune BDFE(O–H) over a broad range, while keeping the same heme-CpdI first-shell sphere of ligands. In the presented models, the $E^{\circ}_{H,CpdI}$ and BDFE(O–H) increase over the range of ca. 10-12 kcal mol⁻¹ as the F_z is altered from –51.4 MV cm⁻¹ to +51.4 MV cm⁻¹ (**Figure 7**), indicating that change of the $E^{\circ}_{H,CpdI}$ of 100 mV is induced by the increase in F_z of ca. 20-24 MV cm⁻¹.

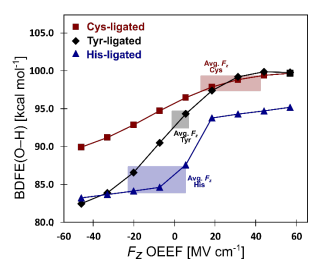


Figure 7. Progression of the BDFE(O–H) in the Cys-, Tyr-, and His-ligated heme-iron models with a change of the F_z component of OEEF. The average values of the LEF F_z observed in the Cys-, Tyr-, and His-ligated proteins from **Table 1**, showing the range of applicable features utilized in Nature are indicated by the faded rectangles. Note that the thermal enthalpic, and entropic contributions to the change of the free energy were neglected (see SI for **Computational Details**).

In more detail, the BDFE(O–H) change with the OEEF is, in fact, a product of the uneven effect of the OEEF on E°_{CpdI} and $pK_{A,CpdII}$ (**Figure 8**); note that BDFE(O–H) = 23.06 x E°_{CpdI} + 1.37 x $pK_{A,CpdII}$ + 57.6 kcal mol⁻¹. In all three models, E°_{CpdI} and $pK_{A,CpdII}$ exhibit the opposite response to the field: growth in F_z increases E°_{CpdI} and decreases $pK_{A,CpdII}$, but $pK_{A,CpdII}$ is approximately half as sensitive. Utilizing Cys-ligated model as an example, we note that as F_z increases by 51.4 MV cm⁻¹, E°_{CpdI} increases by 0.38 V (8.7 kcal mol⁻¹), whereas $pK_{A,CpdII}$ decreases by only 4 units (5.5 kcal mol⁻¹). We expect that this differential tuning of E°_{CpdI} and $pK_{A,CpdII}$ by the LEF could also be utilized by the enzymes, to steer the asynchronicity in the H-atom abstraction between the proton and electron transfers, to achieve a higher reactivity/selectivity.^{13,55}

Finally, we note that the computed properties of all three models (E°_{CpdI} , $E^{\circ}_{H,CpdI}$, and $pK_{A,CpdII}$) respond similarly to positive F_z (Figure 8). However, DFT fails dramatically for E°_{CpdI} and $pK_{A,CpdII}$ in the negative F_z , for the His-ligated model. Since the progression of the $E^{\circ}_{H,CpdI}$ seems to be unaffected by the field orientation, such a failure can be attributed to the incorrect description of the non-protonated Fe^{IV}O CpdII state (*cf.* Scheme 2). Indeed, while the energies of all of the other states from Scheme 2 decrease as the OEEF is applied in either direction (Figure 9), the behavior of His-ligated Fe^{IV}O CpdII diverges, showing the largest energy at F_z of ca. –40 MV cm⁻¹. A deviation from (Trp⁺⁺/Por⁺⁺)Fe^{IV}=O CpdI and Fe^{IV}OH CpdII thus explains a rational behavior of the $E^{\circ}_{H,CpdI}$, but an extraordinary unphysical description of the E°_{CpdI} and $pK_{A,CpdII}$.

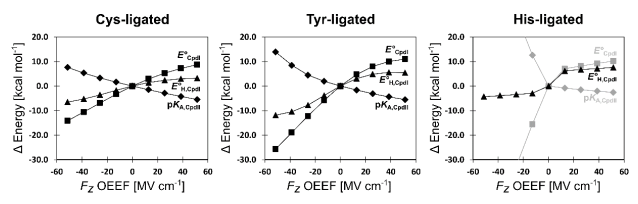


Figure 8. Progression of the physicochemical properties from Scheme 2 of the Cys-, Tyr-, and His-ligated heme-iron models with a change of the F_z component of OEEF. Note that the thermal enthalpic and entropic contributions to the change of the free energy were neglected (see SI for Computational Details).

As analyzed in detail in Figure 9 (see also Supporting Note S1 in the SI), we assign the failure to an increased multi-reference character and/or improper description of the charge delocalization in the Fe^{IV}O CpdII state in the applied field. Although the HOMO (π_{Trp}) and HOMO-1 (a_{1u}) are nearly degenerate in the Fe^{IV}O CpdII in the zero field, OEEF breaks the degeneracy, and influences these orbitals very differently. While the Por a_{1u} and a_{2u} Kohn-Sham orbital energies are only slightly altered by the field in both positive and negative direction (the energy is decreased by ~ 20 kcal mol⁻¹ under OEEF increasing from -51.4 MV cm⁻¹ to +51.4 MV cm⁻¹), the Trp orbital energy changes dramatically, by as much as 103 kcal mol⁻¹. The HOMO character thus inverts from π_{Trp} in the negative fields to Por a_{1u} in the positive fields, and that crossover introduces complications for single-determinant DFT. The result is insensitive to basis set, type of DFT functional, or change in spin state (see **Supporting Note** S1 in the SI). Hence, the asymmetry in the E°_{CpdI} progression at different fields is seen (Figure 8). Finally, to test the hypothesis regarding the multi-reference character, we have evaluated the OEEF effect on the truncated model not including the Trp and Asp residues (Figure S7). This remedied the unphysical behavior in the negative field, but the calculations yield a defective behavior in the positive field, as a result of the incomplete representation of the system. In conclusion, it appears that DFT cannot accurately predict how the thermodynamic properties of the His-ligated peroxidases change with the applied field, but the trends are qualitatively the same for all three model systems. The increase in F_z results in increase of both electron-transfer and proton-coupled electron-transfer potentials, and decrease in the pK_A .

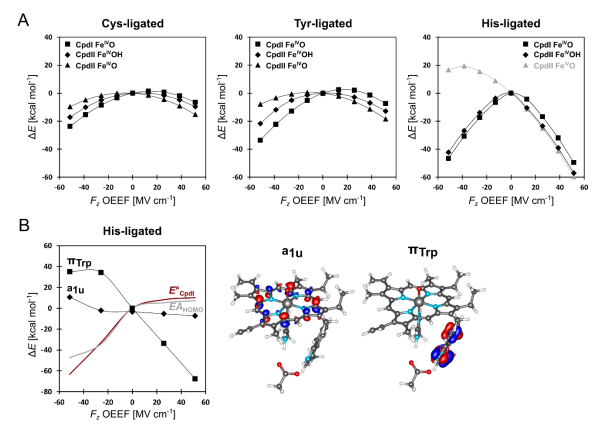


Figure 9. A: Relative change in the energy of the individual states from Scheme 2 with the variation of the OEEF F_z . The CpdI and Fe^{IV}OH CpdII intermediates get stabilized in the positive or negative F_z . In the Hisligated model, the behavior of Fe^{IV}O CpdII deviates significantly from the trend. B: Evolution of the Por a_{1u} and π_{Trp} Kohn-Shamm orbital energies in Fe^{IV}O CpdII with the change of F_z . We note that the change in E°_{CpdI} (*red curve*) calculated directly from the energies of the CpdI and Fe^{IV}O CpdII intermediates corresponds well with the electron affinity (*gray curve*) estimated from the simple change in the HOMO energy of the Fe^{IV}O CpdII. The change in the HOMO character from π_{Trp} to Por a_{1u} is thus suspected for the unphysical calculated electrochemical behavior in negative F_z .

Discussion on the heme-iron proteins selectivity related with the local electric fields in their active sites

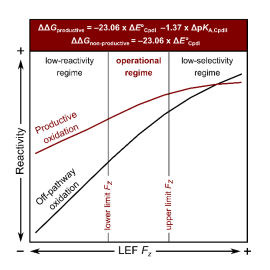
Comparing the large F_z in the P450 monooxygenases with the intermediate or even negative F_z in some catalases and peroxidases (e.g., $\Delta F_z \sim 95$ MV cm⁻¹ between cholesterol 7-alpha-hydroxylase and HRP; *vide supra*) leads us back to a long-standing question: *Why did Nature select a strong electron-donating Cys axial ligand for her most oxidizing heme-iron enzymes*?⁵⁶ We see that LEF in both Cys- vs. Hisligated enzymes can considerably reduce the effect of the axial ligand, bringing the systems closer to oxidative behavior. That is, high F_z in the Cys-ligated cytochrome P450 enzymes (as compared to the lower/negative F_z in peroxidases) increases the electrophilicity of CpdI and decreases the basicity of

CpdII (*cf.* Figure 8), and that eventually "undermines" the effect of the thiolate ligation.⁸ Specifically, the average fields observed in both protein groups (i.e., $F_z = 28.5$ MV cm⁻¹ in Cys-ligated vs. $F_z = -8.7$ MV cm⁻¹ in His-ligated proteins) should be responsible for the shift of ~300-400 mV in E°_{CpdI} and ~3 units in $pK_{A,CpdII}$. Nevertheless, the experimentally obtained values of reduction potentials and basicities of CpdI and CpdII still demonstrate the significantly higher basicity of the Cys-ligated systems. This was envisaged by Green, who proposed a two-fold effect in the P450 catalysis:⁵¹ (*i*) a negative shift of the reduction potential (as compared to the His-ligated system) is compensated by the increase in the pK_A by approximately five units, eventually strengthening the driving force for the H-atom abstraction, *via* a higher contribution of the basicity component to the BDFE(O–H); and (*ii*) a lower reduction potential suppresses the unwanted auto-oxidations of the fragile protein framework. Our results from the previous sections, however, suggest that the situation may become much more complicated in the context of the LEF working unequally on the E°_{CpdI} and $pK_{A,CpdII}$ components of the BDFE(O–H).

Specifically, a finding that BDFE(O–H) is additionally increased in the P450 proteins as the effect of high F_z accords with the high reactivity toward the C–H abstractions. However, we also propose that there must exist both upper and lower limits of F_z for the P450 catalysis to occur, due to the trade-off between productive selectivity and sufficient reactivity (see Scheme 3). Following the work of Green, we suggest the upper limit is represented by the increase of the rate of the non-productive oxidation pathway of the protein framework (for the CYP158 protein studied in ref. ⁵¹, such non-productive pathway corresponds to the oxidation of the adjacent Tyr residues), which is elevated by decreasing the p K_A of the CpdII as F_z increases. A ratio between the non-productive oxidation rate by two different hypothetical CpdI species in otherwise identical systems was derived as: $\frac{k_{z,np}}{k_{n,np}} = 10^{\binom{ApK_A,CpdII}{n}}$, which suggests that an increase in F_z by ca. 17 MV cm⁻¹ should result in an

increase in the rate of the non-productive oxidation by one order of magnitude (*cf.* change in the $pK_{A,CpdII}$ in **Figure 8**, *left*). On the contrary, a lower/negative F_z would lead to a higher ratio between the productive and non-productive reactivity owing to the increase in the CpdII basicity. However, given the uneven effect of the field on the E°_{CpdI} and $pK_{A,CpdII}$, this would simultaneously decrease BDFE(O–H) and, therefore, result in a decreased overall catalytic efficiency. Note that free energy of the reaction is given by: $\Delta G_{\text{productive}} = \text{BDFE}(\text{C}-\text{H}) - \text{BDFE}(\text{O}-\text{H})$; thus, the low reactivity in the productive pathway represents a lower bound of the practicable F_z . All in all, the idea of the confinement of the range of F_z can be supported, e.g., by the positive orientation of the LEF along the

Fe–O axis in 100 % of cases in Cys-ligated proteins (ranging from ca. 3 MV cm⁻¹ to 58 MV cm⁻¹), or by a very narrow range of F_z in the catalases proteins, manifested by a standard deviation of $\sigma(F_z) = 4.7$ MV cm⁻¹.



Scheme 3. A specific range of the LEF F_z is required to be adopted by a protein to accomplish its biological function. The lower and upper bounds of the F_z are given by the counterbalance between the low reactivity on the one side and the low selectivity on the other. Note that reactivity in both productive and non-productive oxidations in the H-atom abstractions is primarily governed by the E°_{CpdI} term, while the steepness with respect to the F_z is decreased in the productive pathway due to the opposite response of the $pK_{A,CpdII}$.

We may also speculate that other steps in the catalytic cycles (even those that are not directly connected with the CpdI or CpdII intermediates) are indirectly linked with the F_z component of the LEF, which may be required to adopt a precisely tuned value for the enzyme canonical function. For example, a decrease in the $E^{\circ}_{H,CpdI}$ and E°_{CpdI} potentials at lower F_z would also likely affect the Fe^{3+/2+} reduction potential of the enzyme's resting state, suppressing the electron delivery from the NAD(P)H, which is a prerequisite for the dioxygen binding. Additionally, as suggested by Winkler and Gray, the failure in the substrate oxidation must be under specific time constraints remedied by an electron transfer to the CpdII to regenerate the secure resting state of the enzyme.^{57–59} Such protection of the protein framework was proposed to be accomplished by the adjacent Tyr or Trp residues, which need to provide reducing equivalents near the CpdII reduction potential. Arguably, the E°_{CpdII} is also tightly coupled with the F_z . Finally, we suggest that proteins' selectivity can be steered by regulating the disparity between E°_{CpdI} and $pK_{A,CpdII}$ via introducing the asynchronicity in the proton/electron transfer in the H-atom abstraction.^{13,55}

CONCLUSIONS

Utilizing a number of computational approaches on the small model systems, it was previously recognized that relatively large oriented electric fields could significantly contribute to the heme-iron reactivity and selectivity. However, a comprehensive association with the field magnitudes inside the proteins' active sites was missing, and the true accessibility of such fields in Nature remained unquestioned. Herein, we bridge the pure computational studies to enzymes, and analyze the electric fields actually present in a wide variety of heme-iron proteins. The local electric fields in proteins appear to be of lower magnitude than studied in small models (the average fields are of ~ 20 MV cm⁻¹), suggesting that the possibility of completely reverting the reaction selectivity (as was done in a small computational model) would be rather difficult. However, we demonstrate that the local electric fields can still significantly contribute to several key mechanistic aspects.

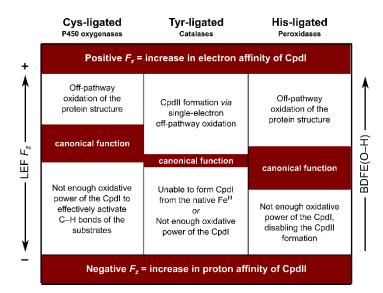
Firstly, the field in enzymes is seen not to be randomly oriented, and indeed exhibit a preferred direction along the Fe–O bond of the supposed CpdI Fe^{IV}O intermediate. This is consistent with the alteration in the oxyl-radical character of the oxo group, modifying the CpdI reactivity/selectivity. The finding suggests, there is indeed a functional role of the field invoked by Nature.

The field of the highest magnitude and preferred orientation along Fe–O is observed in the Cysligated heme-iron proteins, such as P450 oxygenases and peroxygenases. That is consistent with the enhancement of their H-atom-abstraction reactivity as opposed to the strong electron-donation ability of their Cys-axial ligation. We have evaluated that oriented external electric field can increase the bond-dissociation free energy of the Fe^{IV}O–H bond of CpdII (i.e., BDFE(O–H) in **Scheme 2**) by ca. 3 kcal/mol, contributing to the thermodynamic driving force for the H-atom-transfer reactions. Importantly, we noticed that an increase in the BDFE(O–H) is associated with the uneven effect of the field on E°_{CpdI} and p $K_{A,CpdII}$. Consequently, the rise in the H-atom-transfer reactivity is followed by a decrease in the p $K_{A,CpdII}$, concurrently aggravating the chemistry toward the off-pathway oxidation. In detail, the increase in the BDFE(O–H) by 3 kcal/mol is associated with the decrease of p $K_{A,CpdII}$ by 4 units, along with the 100x acceleration of the side redox reactions. We thus suggest there must exist an upper bound of the practicable F_z , given by the increase of the side redox reactivity; and the lower bound F_z , provided by the low overall thermodynamic efficiency (**Scheme 4**; *left*).

We observed fields of intermediate magnitudes within the His-ligated heme-iron proteins, which are additionally oriented either in the positive, or in the negative direction with respect to the Fe–O bond (**Scheme 4**; *right*). While the intermediate magnitudes of the field correspond with the lower reactivity of the CpdI intermediate as compared with the Cys-ligated proteins (i.e., electron

transfer vs. H-atom-transfer reactivity), the greater range of F_z suggests the higher tunability of the thermodynamic properties. In view of the peroxidases subclass of proteins, we successfully correlated the observed local electric fields in the active sites with the experimentally determined CpdI reduction potentials. In addition, we believe that the observed fields in various peroxidases, such as cytochrome *c* peroxidase (C*c*P), *leishmania major* peroxidase (LmP), and ascorbate peroxidase (APX), can bring some understanding into the dilemma of the different resonance forms embraced by these proteins. In Nature, the C*c*P and LmP have unpaired electron located on the adjacent Trp residue and coupled with FeO in Cpd I, whereas APX has a different electronic structure with the unpaired electron located on the porphine ring. We show that the local electric field identified in the peroxidases is consistent with this picture, and is responsible for ~0.5 *e* spin density translocation from Trp to Por in going from C*c*P and LmP to APX.

Finally, the lowest magnitude of the field, reporting no preference for the orientation along Fe–O, was observed in the Tyr-ligated heme-iron catalases. The lower magnitude is again consistent with the lower reactivity of the CpdI in catalases, as compared to cytochromes P450. On the other hand, the narrowest range of the LEF F_z in catalases might arise from the strict requirement of the proteins to hold in check the low-enough E°_{CpdI} so that the single-electron pathway and the formation of the CpdII intermediate is suppressed, and the high-enough reactivity of the Fe³⁺ resting state to alter between the native ferricatalase and CpdI.



Scheme 4. The average F_z is indicative of the CpdI reactivity, i.e., the fact that Cys-ligated proteins report the highest F_z is consistent with their ability to activate strong C–H bonds, as compared to a lower CpdI reactivity in Tyr-ligated and His-ligated oxidoreductases; *cf.* average $F_z = 28.5$ MV cm⁻¹, 3.0 MV cm⁻¹, and -8.7 MV cm⁻¹ for Cys-, Tyr-, and His-ligated proteins. In each of these protein classes, the operational LEF F_z range is given by

the trade-off between the low reactivity of the CpdI intermediate at the more negative F_z and the low selectivity of the reactions at the more positive F_z .

To conclude, although the observed LEFs are not so strong as to divert the reactivity and/or selectivity of the heme-iron enzymes (as was previously inferred from the theoretical investigations of the small model systems) they can significantly contribute to many aspects of the heme-iron chemistry. Hence, LEFs inside the proteins' active sites are an integral part of the metal-oxo reactivity, and as such, they should be included to properly describe the electronic structure, physicochemical properties, and reaction mechanisms of these proteins.

SUPPORTING INFORMATION

The Supporting Information is available free of charge at https://pubs.acs.org//.

Details of the local electric field calculations, proteins preparations, and QM calculations; DFT limitations and multiconfiguration (CASSCF/MC-PDFT) calculations of His-ligated model; Comparison of different DFT approaches; Supporting Figures; Supporting Tables.

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TOC Graphic

