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MD simulations and QM/MM calculations show that single-site mutations of cytochrome P450_{BM3} alter the active site's complexity and the chemoselectivity of oxidation without changing the active species†

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It is a long-standing mechanistic consensus that the mutation of the proton-shuttle mediator Threonine (T) in Cytochrome P450 enzymes severs the water channel and thereby quenches the formation of the active species: the high-valent iron(IV)-oxo porphyrin π -cation radical species, compound I (Cpd I). Using MD simulations and hybrid QM/MM calculations of P450_{BM3} we demonstrate that this is not the case. Thus, while the original water channel is disrupted in the T268A mutant of the enzyme, a new channel is formed that generates Cpd I. With this new understanding, we address the puzzling regiochemical and kinetic-isotope effect (KIE) results (Volz *et al.*, *J. Am. Chem. Soc.*, 2002, 124, 9724–9725) on the sulfoxidation and *N*-dealkylation of dimethyl-(4-methylsulfanyl-phenyl)-amine by wild type (WT) P450_{BM3} and its T268A vs. F87A mutants. We show that the observed variable ratio of *S*/Me oxidation for these enzymes, *vis-à-vis* the constant KIE, originates from Cpd I being the sole oxidant. Thus, while the conserved KIE probes the conserved nature of the transition state, the variable regiochemical *S*/Me ratio reflects the active-site reorganization in the mutants: the shifted location of the new water channel in T268A tightens the binding of the *S*-end by Cpd I and increases the *S*/Me ratio, whereas the absence of π -interaction with the *S*-end in F87A creates a looser binding that lowers the *S*/Me ratio. Our results match the experimental findings. As such, this study sheds light on puzzling experimental results, and may shift a central paradigm in P450 research. The broader implication on enzymatic research is that a single-site mutation is not a localised alteration but one that may lead to a profound change in the active site, sufficiently so as to change the chemoselectivity of catalyzed reactions.

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1. Introduction

The mechanistic elucidation of reactions catalyzed by enzymes often raises enigmas that are inspired by considering simple chemical mechanistic schemes, when in fact these reactions proceed in highly complex and plastic active sites, with multiple residues, water channels, ions, *etc.* Variable chemoselectivity (*e.g.* regioselectivity and enantioselectivity) due to single site mutations in the protein is one such outcome, which often implies that the reactive species of the enzyme has changed its chemical identity/character or reaction mechanism in response

to the mutation. In essence, however, what really changes is the active site, and hence it is essential to also consider the alternative possibility that the single-site mutation induces changes in the complexity of the active site and thereby imparts the observed changes in chemoselectivity. This paper addresses such a general conundrum by focusing on the enzyme P450_{BM3}, which is a member of the superfamily of cytochrome P450 enzymes.^{1–9}

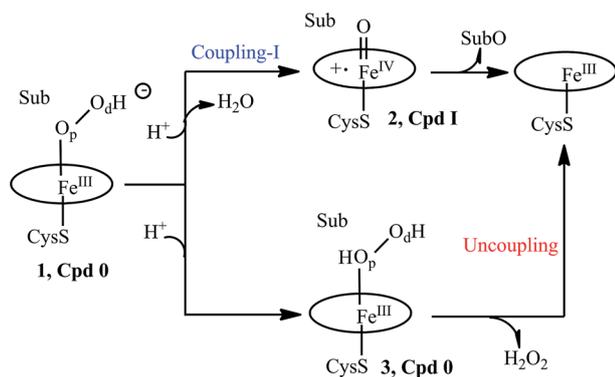
It is generally thought that P450 enzymes use multiple oxidant species^{1,2,6,7,10} because single site mutations, which sever the proton shuttle pathways required to generate the principal oxidant species of the enzyme (2 in Scheme 1), cause a change in the regioselectivity of oxidation. One of these long-standing puzzles is the variable regioselectivity of sulfoxidation vs. *N*-dealkylation observed during the oxidation of dimethyl-(4-methylsulfanyl-phenyl) amine (**Sub1** in Scheme 2), by wild-type (WT) P450_{BM3} and its T268A and F87A single-site mutants (MTs).⁶ Using molecular dynamics (MD) with sufficiently long timescales and hybrid QM/MM calculations, we address here the WT and the two MTs, as well as the variable oxidative

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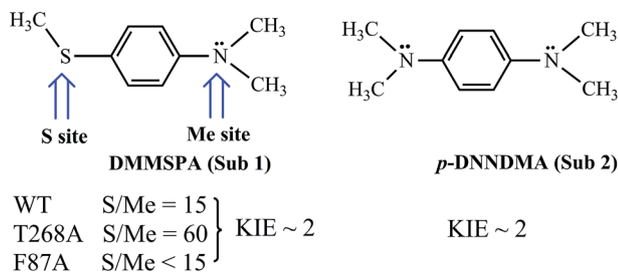
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Scheme 1 A truncated catalytic cycle of P450.

Scheme 2 A summary of the regioselectivity and kinetic isotope effect (KIE) data concerning the substrates DMMSPA (Sub 1) and p-DNNDMA (Sub 2) by wild-type (WT) P450_{BM3} and its T268A and F87A mutants (MTs).

regioselectivity and the constant KIE. As the story unfolds, our study will demonstrate that the selectivity alterations reflect major reorganization in the active sites, *e.g.*, the formation of a new water channel,¹¹ that affect the substrate juxtaposition *vis-à-vis* the active species of the enzyme. Single-site mutations are not merely localised alterations.

2. The mechanistic dilemma and chemical puzzles in P450_{BM3} and some of its mutants

Let us first present the mechanistic dilemma, and then proceed with the experimental background for invoking the two-oxidant scenario by Volz *et al.*⁶ A truncated, generic catalytic cycle of P450 is shown in Scheme 1, starting from the PorFe^{III}-OOH⁻ species called compound 0 (Cpd 0), **1**. Protonation of **1** on the distal oxygen (O_d) leads to the liberation of a water molecule and the formation of the ultimate oxidant, **2**, the porphyrin π -cation radical ferryl-species (Por⁺Fe^{IV}=O), so-called Cpd I. Cpd I leads to the oxidation of the substrate (Sub), and is the consensus oxidant, responsible for the vast majority of the P450 catalyzed mono-oxygenation reactions.¹⁻¹⁵ This process is referred⁷ to as “Coupling-I” because it couples the O₂ activation to the substrate oxidation. This is also the native process of P450s.

However, a single mutation of the Thr residue to Ala (T \rightarrow A) is thought to disrupt the proton-shuttle machinery of the enzyme and increase the amount/lifetime of Cpd 0 (ref. 7, 10, 11 and 16–19) that now undergoes protonation on the proximal oxygen (O_p) of **1**, leading to the formation of the ferric hydrogen peroxide complex, PorFe^{III}(O₂H₂) **3**. Since the Fe–O bond in **3** is rather weak, much depends on the persistence of **3**. Usually, **3** liberates hydrogen peroxide in a process called “uncoupling” which is an aborted oxidation. This leaves Cpd 0 (**1**) as a possible oxidant in the T \rightarrow A mutants of P450. Therefore, the consideration of Cpd I and Cpd 0 as the potential oxidants in P450s and their T \rightarrow A mutants (MTs) has become a mechanistic consensus in the field.

Let us turn now to Scheme 2 and inspect the intriguing mechanistic results of Volz *et al.*,⁶ which imply the presence of Cpd 0 alongside Cpd I. Thus, the WT enzyme sulfoxidates the sulfur site of Sub 1 15 fold more than dealkylating the NMe₂ site *via* the initial C–H hydroxylation. The MT T268A – thought to quench Cpd I formation and use Cpd 0 as an oxidant by the existing mechanistic paradigm – was found⁶ to increase the S/Me ratio to 60, whereas the MT F87A that is not thought to affect the proton-shuttle machinery actually decreases the S/Me ratio to below 15.

The mutations and changes in the S/Me reactivity-ratio do not affect, however, the intrinsic kinetic isotope effect KIE_{H/D} (~2), which is determined for the appropriately deuterated Sub 1. Furthermore, to rule out the slow interchange of the two sites of Sub 1, Volz *et al.* employed an analogous substrate Sub 2, and observed that both the NMe₂ and NMe₂[D-6] (D-6 indicates the number of substituted deuterium atoms at the two methyl groups) sites of the latter undergo oxidation with a KIE_{H/D} of ~2, for both the WT and MT T268A enzymes. As such, all of these findings, taken together, were cautiously interpreted⁶ in a scenario whereby Cpd I and Cpd 0 co-exist in both the WT and the MT T268A enzymes, where Cpd I performs N-dealkylation whereas Cpd 0 performs sulfoxidation. As such, the S/Me ratio has to reflect the relative abundance of the two oxidants.

Similarly, Cryle and de Voss^{20a} investigated the oxidation of fatty acids and the ω -2-thia-substituted ones by P450_{BM3} WT and MT T268A enzymes, and showed that C–H hydroxylation suffered high uncoupling in the MT T268A enzyme, whereas sulfoxidation was coupled in both the MT and WT enzymes. Nevertheless, both reactions conserved similar regio- and enantio-selectivities in the MT and WT enzymes. Thus, once again, this evidence seems to compellingly suggest that in both the WT and MT enzymes the two oxidants function in tandem; Cpd I is responsible for C–H hydroxylation, while Cpd 0 carries out the sulfoxidation.

We note that besides the two-oxidant scenario, Volz *et al.*⁶ considered that the same result could have been caused by changes in the structure of the active site due to mutations. Moreover, Cryle and de Voss^{20b} showed more recently that the observed uncoupling – due to the presumed “loss” of Cpd I’s activity in favor of Cpd 0’s – was highly substrate dependent, while the information obtained from the radical clocks during C–H hydroxylation supported Cpd I as a single oxidant.²¹ Additionally, all known computational results for heme systems



show that Cpd 0 is a very poor oxidant compared to Cpd I^{22,23} and it has extremely high sulfoxidation barriers.²³ As such, Cpd 0 cannot really be responsible for the sulfoxidation of **Sub1** in the presence of Cpd I. In fact, we⁷ and Altarsha *et al.*²⁴ showed that in the MT T252A of P450_{cam}, Cpd 0 is protonated very easily to form PorFe^{III}(O₂H₂), **3**, and will not survive to carry out substrate oxidation.

In view of all of the intriguing results, Cpd I may well be the sole oxidant responsible for oxidation in the WT enzyme *via* the native coupling-I process. In the MT T268A of P450_{BM3}, the formation of Cpd I may either occur *via* the coupling-I process or the recently found⁷ coupling-II pathway. However, the latter process requires a sufficiently large and tightly bound substrate that can barricade the H₂O₂ moiety.⁷ Otherwise, PorFe^{III}(O₂H₂) will lose H₂O₂, thus resulting in uncoupling (Scheme 1). As mentioned already, our results show that, in P450_{BM3}, PorFe^{III}(O₂H₂) is unstable with respect to a loss of H₂O₂ (see ESI, Fig. S1†). We are therefore left with Cpd I as the only possible oxidant. But how does Cpd I actually form and how does it concoct the intriguing scenario described by Volz *et al.*?⁶ The present study answers these questions by means of extensive MD simulation sets of 100–200 ns of the species, combined with QM/MM calculations.²⁵ First, we address the mechanisms of Cpd I formation in the T268A mutant of P450_{BM3}, and then we explore the mechanism of **Sub1** oxidation by P450_{BM3} and its mutants. Our findings will show that changes are required in some key mechanistic paradigms in P450 mechanistic research.

3. Computational methods and details

3.1. System preparation and setup

The substrate binding site in P450_{BM3} is well known, and therefore we started with a palmitic acid-bound P450_{BM3} (PDB no. 1FAG)⁸ and replaced the palmitic acid with **Sub1** to use an experimentally verified binding site. Since the crystal structure of P450_{BM3} with **Sub1** is not available, we used two different initial orientations of **Sub1** to confirm the proper binding of the substrate; one where the N-(CH₃)₂ group is pointing towards the Fe=O end and the other where the S-CH₃ group points towards Fe=O in the WT and T268A mutant.

Missing hydrogen atoms were added by the LEaP module of Amber 14.²⁶ The study considered three of the P450_{BM3} species of interest (Cpd 0, Cpd I and Fe(H₂O₂)). The force field for Cpd I was taken from the literature,²⁷ while the force fields for the Cpd 0 and Fe(H₂O₂) states were parameterized using “MCPB.py” modeling.²⁸ The partial atomic charges and missing parameters for **Sub1** were obtained by the RESP (restrained electrostatic potential) method,^{29,30} using HF/6-31G* calculations. A few Na⁺ ions were added to the protein surface to neutralize the total charge of the systems. Finally, the resulting system was solvated in a rectangular box of TIP3P³¹ water extending up to a minimum cutoff of 10 Å from the protein boundary. The Amber ff14SB force field was employed for the protein in all of the MD simulations.

3.2. MD simulations

After proper parameterizations and setup, the resulting system's geometries were minimized (5000 steps for the steepest descent and 10 000 steps for the conjugate gradient) to remove the poor contacts and relax the system. The systems were then gently annealed from 10 to 300 K under a canonical ensemble for 50 ps with a weak restraint of 5 kcal mol⁻¹ Å⁻². Subsequently, the systems were maintained for 1 ns of density equilibration under an isothermal–isobaric ensemble at a target temperature of 300 K and a target pressure of 1.0 atm using a Langevin-thermostat³² and a Berendsen barostat³³ with a collision frequency of 2 ps and a pressure-relaxation time of 1 ps with a weak restraint of 1 kcal mol⁻¹ Å⁻². This 1 ns of density equilibration is not identical with conformational equilibration, but is rather a weakly restrained MD in which one slowly relaxes the system to achieve a uniform density after heating dynamically under periodic boundary conditions. Thereafter, we removed all of the restraints applied during the heating and density dynamics and further equilibrated the systems for ~3 ns to get well settled pressure and temperature for conformational and chemical analyses.

Finally, a productive MD run was conducted for each system for at least 100 ns. In general, 100 ns of MD sampling is sufficient for ligand binding and substrate recognition. In some cases, in order to ascertain the convergence of the MD, we proceeded with an additional 100 ns simulation, which did not change the substrate orientation that was obtained in the initial 100 ns. As such, we consider that this convergence test is sufficient for sampling the binding of **Sub1** in the present study. The most populated structures were calculated by the clustering of MD trajectories by the CPPTRAJ module of Amber 14. During all of the MD simulations, the covalent bonds containing hydrogen were constrained using SHAKE,³⁴ and particle mesh Ewald (PME)³⁵ was used to treat the long-range electrostatic interactions. We used an integration step of 2 fs during the entire simulation. All of the MD simulations were performed with the GPU version of the Amber 14 package.³⁶

3.3. QM/MM methodology

For the subsequent QM/MM calculations,³⁷ we used representative snapshots extracted from the most populated structures within the clustering of the MD trajectories. This choice is statistically more rigorous than stochastically chosen snapshots. All of the water molecules beyond the sphere of the enzyme were removed, and the resulting enzyme was solvated with a 16 Å layer of TIP3P water molecules (yielding a total of *ca.* 45 600 atoms).

The propionate-truncated heme and the entire **Sub1** were included in the QM region, whereas all of the protein residues and water within 8 Å of the heme and **Sub1** were included in the active region. The atoms in the active region interact with the QM atoms through electrostatic and van der Waals interactions and were considered in the subsequent QM/MM calculations. As such, the QM/MM results also account for the interactions of the QM system with key residues *e.g.* Phe 87, Thr 268 and other water residues within 8 Å of the heme and **Sub1**. All of the QM/



MM calculations were performed using ChemShell,³⁸ by combining Turbomole³⁹ for the QM part and DL_POLY⁴⁰ for the MM part. The CHARMM27 force field was employed for the MM region. The electronic embedding scheme⁴¹ was used to account for the polarizing effect of the enzyme environment on the QM region. Hydrogen link atoms with the charge-shift model^{35,37a} were applied to treat the QM/MM boundary.

During the QM/MM geometry optimizations, the QM region was treated by the hybrid UB3LYP⁴² functional with two basis sets. For the geometry optimization and frequency calculation, we used the polarized double- ζ basis set LACVP for iron and the 6-31G(d) basis set for all of the other atoms, collectively labeled as B1. The energies are further corrected with the large all-electron basis-set Def2-TZVP, labeled as B2. All of the QM/MM transition states (TSS) were located by relaxed potential energy surface (PES) scans followed by full TS optimizations using the P-RFO optimizer implemented in HDLC code⁴³ and DI-find.⁴⁴ The QM/MM calculations used the hybrid UB3LYP functional. For Cpd I mediated reactions, the $S = 3/2$ and $S = 1/2$ states exhibit generally similar reactivities.^{7,45} As such, all of the reactions of Cpd I were studied in the $S = 1/2$ state.

4. Results and discussion

Let us start with the mechanisms of Cpd I formation and then proceed to the mechanisms of the oxidation of **Sub 1** in the WT enzyme, and in the T268 and F87A mutants.

4.1. Cpd I formation in the T268A mutant of P450_{BM3}

The anticipation of a second oxidant is based on the conjecture that the T268A mutant quenches Cpd I formation or decreases its production. In fact, we show here that T268A generates Cpd I efficiently *via* the appearance of a very intriguing new water channel due to the mutation. We first ran an MD simulation of 100 ns on Cpd 0 (see ESI, page S2, Fig. S1 and S2†), from which we took two snapshots from the two conformational basins we identified, and performed QM/MM calculations. Fig. 1a shows the results for the snapshot from the major conformer. It is seen that residue Glu267 is nicely connected to the O_d of Cpd 0 through a few water molecules. The proton transfer from Glu267 to O_d generates Cpd I in a single step, with a rather small barrier of 11.3 kcal mol⁻¹. In the minor conformer (Fig. 1b), the channel leads to the protonation of the distal O of Cpd 0 and forms Fe(H₂O₂). To rule out the possibility that the coupling-II⁷ pathway operates in P450_{BM3}, we ran 100 ns MD simulations on the as-formed Fe(H₂O₂). The simulations indicate that **Sub1** is quite flexible in the pocket and is far away from Fe(H₂O₂) (see ESI, Fig. S3†). This in turn suggests that **Sub1** is unable to barricade the H₂O₂ ligand so that Fe(H₂O₂) would lose H₂O₂ resulting in uncoupling (Scheme 1). Such a scenario seems to be consistent with experiments that show that the T268A mutant of BM3 always leads to a partial uncoupling in the presence of fatty acid substrates.^{11,20}

4.2. MD simulation results as the preparation for **Sub1** oxidation

To test the regiochemical preference of the binding sites of the S or N of **Sub1** by Cpd I, we carried out MD simulations of the Cpd I/**Sub1** complex within the active site. We used two different initial orientations of **Sub1** called pose-1 and pose-2; pose-1 refers to the conformation where the N-(CH₃)₂ group points towards the Fe=O end and pose-2 refers to the one where the S-CH₃ group points towards Fe=O. Starting the MD simulations from pose-1, **Sub1** ended up moving away and remained far from the Fe=O moiety during the entire simulation time. This was judged by the distance of the closest N-CH₃ terminal of **Sub1** to Fe=O, which was >10 Å during the entire simulations (see Fig. S4 and Video VS1 in ESI†). To double-check this conclusion, we used molecular mechanical Poisson Boltzmann surface area (MMPBSA) calculations, and calculated the enthalpy changes ($\Delta H = \Delta H_{\text{complex}} - [\Delta H_{\text{protein}} + \Delta H_{\text{Sub1}}]$) upon the binding of **Sub1** for these two poses. The MMPBSA calculations also showed that the binding of **Sub1** in pose-1 is significantly weaker than that in pose-2 (see Table S2† for details). Thus, both the MD simulations and MMPBSA calculations show that pose-1 does not represent an appropriate model for the experimental observations. As such, this orientation was dropped, and the MD simulations were done from pose-2 of **Sub1**, and are presented in Fig. 2.

Fig. 2a shows the fluctuations of the oxidizable moieties of **Sub1** (H in N-CH₃ and S in S-CH₃) relative to Cpd I's oxo atom. It is seen that the substrate is initially far away from Cpd I (>10 Å), but within 5 ns, **Sub1** slides closer to Cpd I and achieves proper juxtapositions *vis-à-vis* the Fe=O moiety. Looking at the distance *vs.* time plot in Fig. 2a, the 90–100 ns interval of the MD simulations shows that the NMe₂ group, which was up to this point far away, now approached the Fe=O moiety. This may raise the possibility that the NMe₂ group may permanently settle close to the Fe=O moiety, even at later times in the simulations. To test this possibility, we extended the MD simulations to 200 ns to check the persistence of the proximity of NMe₂ to the Fe=O end. The extended MD simulation in Fig. 2a shows that it is the sulfur terminal that remains closest to Cpd I for most of the 200 ns, while the NMe₂ group is quite far away for most of the simulation time, with the exception of the 90–100 ns interval. To further characterize the substrate's orientational distribution, we performed the clustering of the MD trajectories for the entire 200 ns of simulation into the related conformational basins (see Fig. S5,† page S5, for three typical snapshots during clustering). This enabled us to verify that in all of the clusters of the MD trajectory, the sulfur end is indeed closer to Cpd I and this is the most populated conformation, in agreement with Fig. 2a.

The reason for the closer binding of the S site of **Sub1** to Cpd I was found to be the interaction of S with the Phe87 residue. The noncovalent interaction⁴⁶ of Phe87 with **Sub1** is shown in Fig. 2b for the initial snapshot and one taken from the 80–100 ns interval; the green patch signifies the S/ π interaction. Judging from the intensity and area of these green patches, it is clear from the two snapshots that the phenyl ring of Phe87



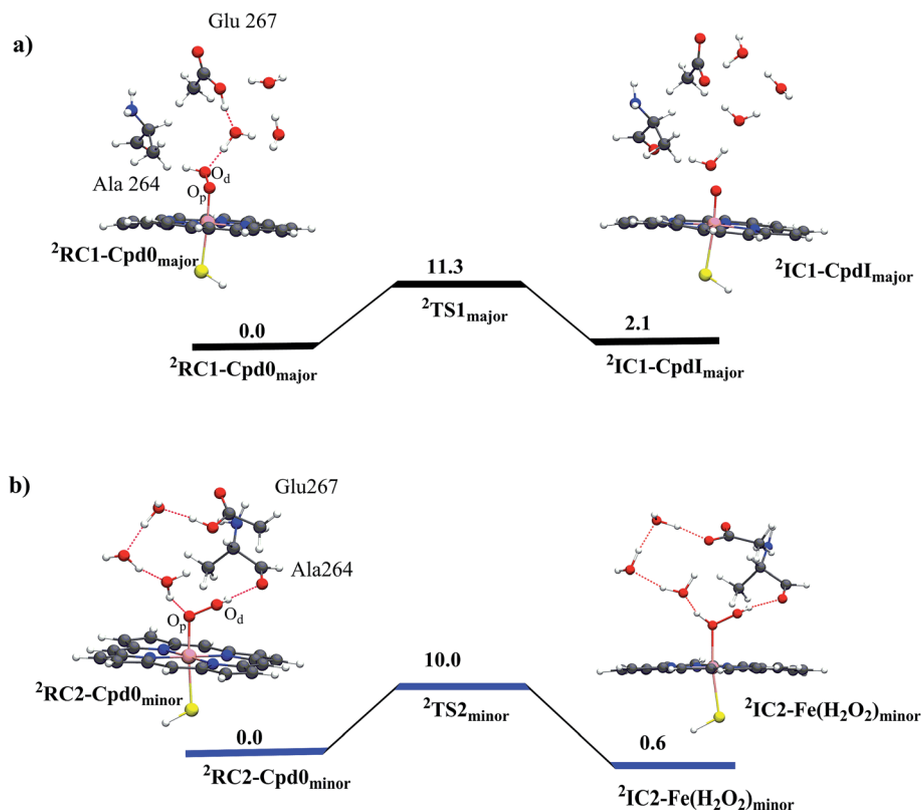


Fig. 1 QM/MM (UB3LYP/B2//B1) energy profiles (in kcal mol⁻¹) for Cpd I formation from Cpd 0 in the T268A mutant: (a) the coupling-I pathway for the major conformer leads to Cpd I formation and (b) protonation of Cpd 0 in the minor conformer leads to Fe(H₂O₂) formation. The energy barriers include ZPE corrections. **Sub 1** is included in the MM part, and MD simulations show that it is located too far from Fe(H₂O₂) to barricade H₂O₂ against detachment (ESI, Fig. S3†). Thus, Cpd I formation from Fe(H₂O₂) (coupling-II⁷) can be ruled out in the T268A mutant of BM3.

intensifies its pull of the S-CH₃ terminal of **Sub1**, which gets closer to Phe87 *via* this interaction. As such, the sulfur terminal is pulled away from Fe=O while NMe₂ slides momentarily closer to the Fe=O end. This rearrangement creates a minor conformational basin in which NMe₂ assumes an oxidizable distance to Fe=O. After 100 ns or so, S restores its proximity to Fe=O, such that the major conformational basin maintains a closer sulfur/Cpd I distance. As such, if the juxtaposition determines the fate of the complex, our MD results predict that sulfoxidation would lead to the major product while C-H activation (of the N-CH₃ moiety) would lead to the minor one, unless the relative barriers predict otherwise. Furthermore, the orientation towards sulfur is preferentially S-prochiral, so provided the MD determines enantioselection the reaction should result in the S-enantioselective sulfoxide.

4.3. QM/MM results of the WT enzyme

To test these MD-based predictions, we performed QM/MM calculations on representative structures of two differently populated conformational basins; one corresponds to the major basin where the sulfur is closest to Cpd I, the other to a minority basin where the C-H bond is the closer moiety to Cpd I (Fig. S5a and c†). Fig. 3 displays the two reactant-complexes, ${}^2\text{RC}(\text{major})$ and ${}^2\text{RC}(\text{minor})$, corresponding to these basins. Analysis of the MD trajectories shows that the frequencies of visits in the two

basins are 1866 (major) vs. 11 (minor). These relative-visit-frequencies plugged into the Boltzmann equation yield the relative energy of the RCs, which is 2.8 kcal mol⁻¹ in favor of the major conformation at the experimental temperature of 303 K. The energy profiles for sulfoxidation and C-H activation, nascent from these RCs, are shown in Fig. 3, along with the structures of the lowest lying transition states for hydroxylation and sulfoxidation, where the energies are given relative to ${}^2\text{RC}(\text{major})$.

Relative to ${}^2\text{RC}(\text{major})$ as the reference energy, the lowest QM/MM barriers (with ZPE corrections) are 10.1 kcal mol⁻¹ for H-abstraction through the minor channel and 9.4 kcal mol⁻¹ for sulfoxidation *via* the major channel. Using the barrier energy difference, 0.7 kcal mol⁻¹, the predicted S/Me ratio at 303 K is 4 : 1, compared with 15 : 1 as reported by Volz *et al.*⁶ (corresponding to a 1.5 kcal mol⁻¹ energy difference). Using another snapshot (see ESI, Fig. S6†), the corresponding barriers lead to the same conclusion that sulfoxidation is preferred over H-abstraction. This result is in qualitative agreement with experimental observations that sulfoxidation would lead to the major and C-H hydroxylation to the minor product in the WT enzyme.⁶

Looking at the other two barriers in Fig. 3, we can see that the reactions from the poorly juxtaposed RCs have high barriers. Thus, the sulfoxidation barrier in the minor basin is 14.6 kcal



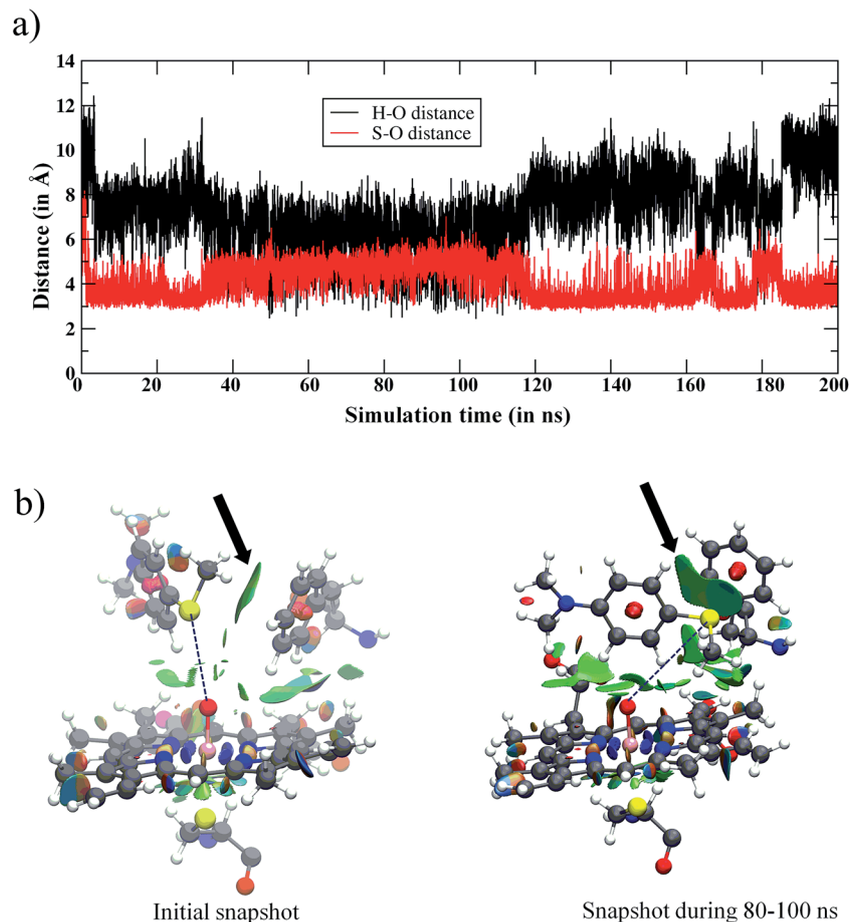


Fig. 2 (a) The variation of the Fe=O \cdots H and Fe=O \cdots S distances vs. simulation time for the WT enzyme. Initially, the amine and sulfur ends of **Sub1** are far away from iron-oxo, however, in the later stages both ends become accessible to Cpd I. However, the major basin of the snapshots places the sulfur end closest to Cpd I. (b) The structures in the initial snapshot and the one during the 80–100 ns of the MD trajectory are shown along with the noncovalent interactions of **Sub1** with Phe87 and Cpd I. The green surfaces show the favorable noncovalent interactions and the area of the green patches indicates the interactions' intensity. Note the increase in the intensity between the sulfur atom of **Sub1** and Phe87 (highlighted by arrows) from the initial trajectory to the one in the 80–100 ns interval.

mol⁻¹, whereas the H-abstraction barrier in the major basin, which highly disfavors N-CH₃-juxtaposition, is huge at 26.5 kcal mol⁻¹. These are much higher barriers than the ones discussed above and as such these pathways will clearly not contribute to the reaction.

To further articulate the reasons for these high barriers, we investigated the charge and spin densities of the species in the minor and major basins (see Fig. 3 and Table S1†). As seen from the spin density (ρ) on the substrate in ²RC(minor), there is a transfer of one electron from the substrate to the porphyrin ring of Cpd I, so that ²RC(minor) involves a single electron reduced Cpd I (so-called Cpd II) with **Sub1**⁺ that diminishes the reactivity in the minor basin, relative to ²RC(major) which involves a cluster of Cpd I with **Sub1**. Indeed, due to the decreased reactivity of Cpd II, the electronic contribution (E_{QM}) to the energy barrier for the H-abstraction for ²RC(minor) (16.9 kcal mol⁻¹) is higher than the electronic contribution to the energy barrier for ²RC(major) (11.2 kcal mol⁻¹).

4.4. MD and QM/MM results of the T268A MT enzyme

The enhanced S/Me regioselectivity of the T268A MT enzyme⁶ with **Sub1** raises an obvious question: how can a single mutation change the regio-selectivity of substrate oxidation? As written above, past explanations of this change^{6,20a} assumed that Cpd 0 is present alongside Cpd I, and the fraction or lifetime of Cpd 0 increases in the T268A MT. Furthermore, it was assumed that Cpd 0 is the species responsible for sulfoxidation, whereas Cpd I is responsible for hydroxylation. However, since all computations show that Cpd 0 has high substrate-oxidation barriers,^{22,23} the assumption that Cpd 0 is responsible for the S/Me-regioselectivity enhancement is deemed questionable. Therefore, we performed MD simulations for the T268A MT to study the substrate reorientation caused by the mutation. As expected, this MD simulation shows completely different behavior to the WT simulation. Thus, in contrast to the WT enzyme, which showed the dynamic behavior of **Sub1** (Fig. 2), in the MT **Sub1** adopts a stable orientation. And as seen in Fig. 4, now the sulfur end is always close to the oxo end of Cpd I, while



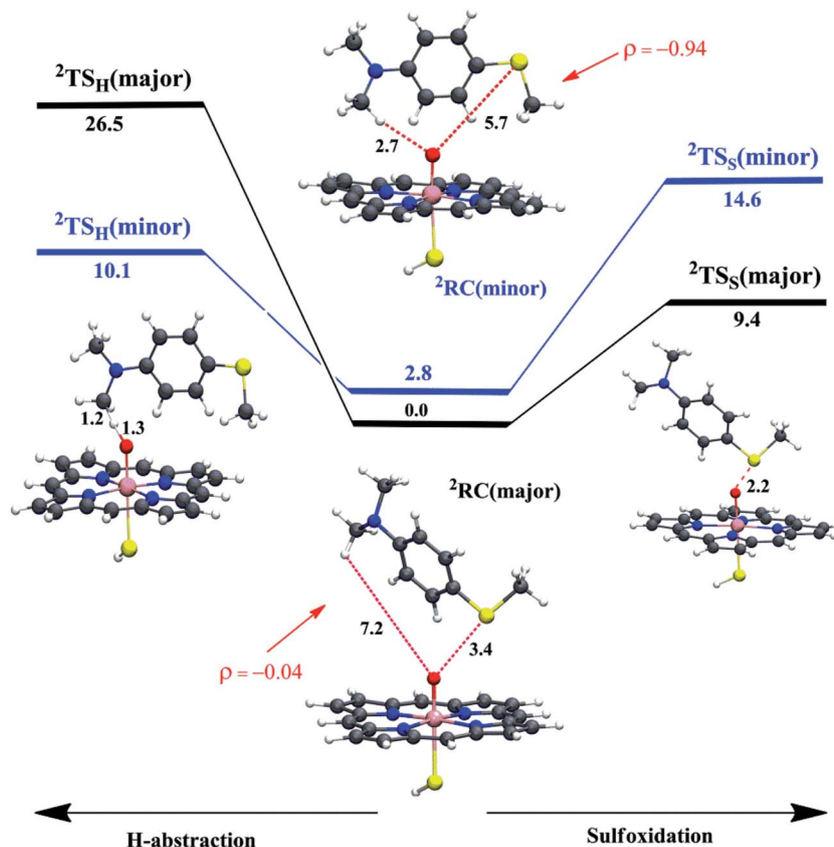


Fig. 3 The QM/MM (UB3LYP/B2//B1) energy profiles (in kcal mol⁻¹) for WT P450_{BM3} (H-abstraction is to the left and sulfoxidation is to the right) nascent from the major and minor reactant-clusters (²RC(major) and ²RC(minor)) and going to the corresponding ²TS_H and ²TS_S. The energy barriers include ZPE corrections. The spin densities (ρ) on the substrate indicate that the oxidant in ²RC(major) corresponds to Cpd I while in ²RC(minor) it corresponds to Cpd II (see also Table S1†, page S5). Only the lowest TS structures are shown (the higher ones can be found in the ESI†), along with key distances (in Å).

the NMe₂ end seldom approaches Cpd I. This is also well supported by the clustering of the trajectories, wherein no cluster represents an NMe₂ end being closer to Cpd I than the sulfur end (see Fig. S7†). This clearly shows that the T268A mutation brings the sulfur end closer to Cpd I and pushes the NMe₂ end away, hence causing enhanced sulfoxidation over H-abstraction.

Since, by analogy to T252 in P450_{cam},¹⁷ the primary role of T268 in P450_{BM3} is thought to be proton delivery *via* an organized water channel, we carefully monitored the water channel and permeability for the entire MD trajectories of the WT and T268A MT enzymes. Identical numbers of permeated water molecules were found for the WT and T268A MT enzymes (See ESI for details,† page S7). However, what completely changed for the T268A MT is the location of the water channel. As shown

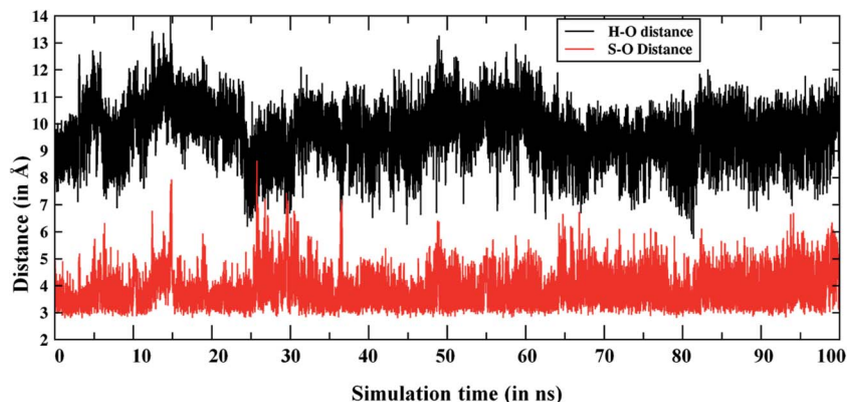


Fig. 4 Comparison of Fe=O...S and Fe=O...H [the H in N(CH₃)₂] distances during the MD simulations of the T268A mutant.



in Fig. 5a, throughout the MD simulations of the WT enzyme the Thr268 residue mediates a very organized water chain (see ESI videos VS2 for the WT and VS3 for the T268A mutant†). The T268A mutation disrupts this channel and instead provides a new aqueduct where the substrate acts as a “wall” to direct the water flow. As a result, the substrate is reoriented and the S terminal gets closer to Cpd I. As shown in Fig. 5b (left item), the water molecules of the new channel, jointly with channel residues, barricade the substrate and fix the S-end close to Cpd I, thus enhancing sulfoxidation. Indeed, as seen from the energy profiles in Fig. 5b (right item), the sulfoxidation barrier is lowered (compared to Fig. 2 for the WT), while the H-abstraction barrier is raised, which is in qualitative agreement with experimental data.⁶ The unique role of the substrate is in line with the observations of Cryle and de Voss.^{20b} Furthermore, here we can see how a major mechanistic paradigm for P450 must be revised in view of these findings. Thus, the T268A mutation in P450_{BM3} destroys the “old” proton shuttle pathway and recreates a new one, which leads to the formation of Cpd I (Fig. 1a). Meanwhile, because the new water channel is located to the far right of the WT’s channel (Fig. 5a, right-hand drawing), the T268A mutation changes the reorientation of the substrate and thus impacts the regioselectivity of the oxidation of **Sub1** by tightly juxtaposing the S end towards Cpd I.

4.5. MD results for the F87A MT enzyme

As discussed above, the F87A mutation decreases the amount of the sulfoxidation product.⁶ Therefore, we also performed MD simulations for the F87A MT. As expected, Fig. 6 shows that now

the S-terminal in the F87A MT (red color) is farther away from the oxo ligand of Cpd I, in comparison to that in the T268A MT (and also in the WT, see Fig. 2a). As already discussed, Phe87 forms a good S/ π interaction that is crucial for the substrate orientation (Fig. 2b). Therefore, mutating Phe87 to Ala eliminates this interaction and causes a different orientation of the S-terminal in the substrate, which might be responsible for the decreased sulfoxidation as seen in experiments.

A closer inspection of the MD trajectories for the F87A mutant (see ESI video VS4†) demonstrates that the routes for the water channel are similar to those for the WT enzyme, but different compared to those found for the T268A MT. Clearly, the T268A MT has a profound impact on the regioselectivity and reactivity of P450_{BM3}. The T268A MT completely alters the water channel and its location in the active site. With the new channel, there is also a new mechanism for Cpd I formation (Fig. 1). Additionally, the shifted location of the water channel alters the binding conformation of the substrate (Fig. 5a and b), and thereby increases the S/Me regioselectivity ratio. On the other hand, F87A conserves the water pathways as in the WT, but it lacks the π /S glue to bring the S terminal as close to Cpd I, thus lowering the S/Me regioselectivity. Our findings strongly support the notion that what causes the changes in regioselectivity are the modifications of the active site in response to the specific mutations. Meanwhile, Cpd I is conserved in the three enzymes, and its S/Me regioselectivity is determined by the modifications in the substrate binding exerted by the T268A and F87A mutations. We further verified that the KIE values in WT and MT T268A are both close to ~ 2 (see ESI,† Page S8) and reflect the constant nature of the H-abstraction TS of the Cpd I species.

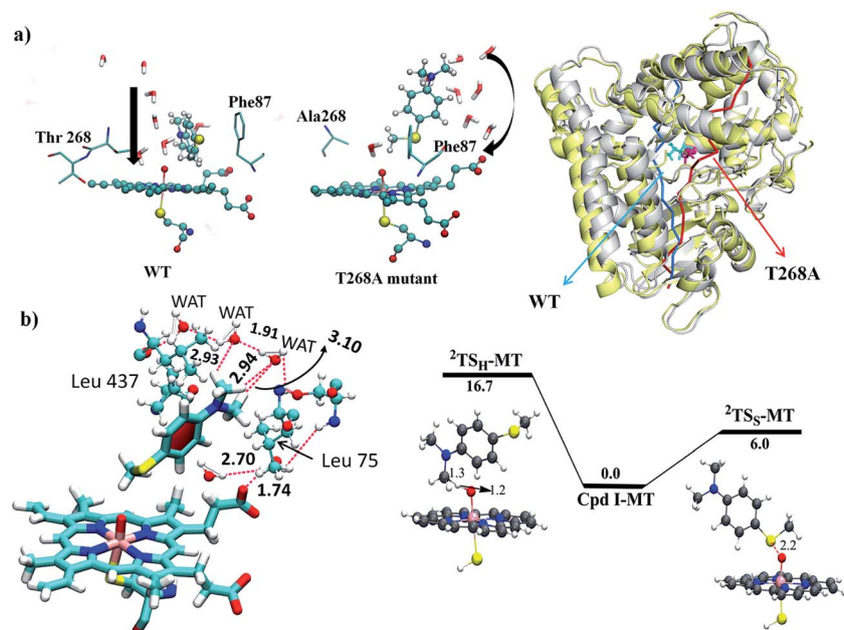


Fig. 5 (a) The water permeation routes were calculated using Pore Walker and are shown near Cpd I in the WT and the T268A MT enzymes. On the right side the two enzymes (the WT is in yellow and the MT is in grey) are overlapped and their water routes are drawn by color-coded arrows; a bluish arrow for the WT and a reddish arrow for the MT. Note that the substrate is acting as a guiding wall for the water stream in T268A. (b) The left item shows water molecules jointly with channel residues barricading the substrate and fixing its S-end near Cpd I. The right hand drawing shows the QM/MM energies and TS structures for H-abstraction (${}^2\text{TS}_\text{H-MT}$) and sulfoxidation (${}^2\text{TS}_\text{S-MT}$).



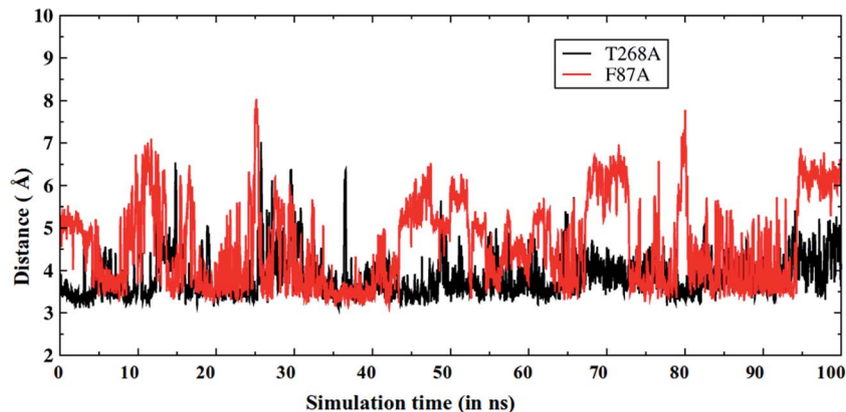


Fig. 6 The Fe=O...S distance vs. the MD simulation time for T268A (black) and F87A (red) mutants. Note that the Fe=O...S distance is longer in the F87A MT enzyme.

5. Conclusions

In conclusion, all of the above findings are consistent with a scenario whereby sulfoxidation and hydroxylation are both mediated by the active species Cpd I, in the WT as well as the MT enzymes. The T → A mutation of P450_{BM3} generates a newly located water channel, guided by Glu267, thus highlighting the flexibility of the enzyme active sites in enzymatic catalysis.^{9,47} This new water channel forms Cpd I very efficiently (Fig. 1a) but this is in competition with uncoupling to H₂O₂ (Fig. 1b) which is consistent with experimental observations.^{11,20b} The variable S/Me ratio originates from variations of the active site in response to the mutations. Thus, since Phe87 confers a π /S interaction that brings the S terminal of the substrate close to Cpd I in the WT enzyme, the lack of Phe87 due to the F87A mutation is a reason behind the decreased yield of sulfoxidation. In the T268A MT, the S/Cpd I proximity increases due to the relocated water channel, and the role of the substrate as a wall for the aqueduct. In contrast, the constant KIE (see ESI,† Page S8) is in line with a “conserved” transition state due to the presence of Cpd I as the sole oxidant in both the WT and T268A.

The very efficient formation of Cpd I in the T268 mutant, along with the mutation-induced active site reorganization that causes an altered water route and an altered orientation of substrate, provides a new paradigm for understanding the regio- and stereo-selectivity issues in Cpd I-mediated reactions. Finally, the study leads to an additional modification of a tacitly assumed mechanistic paradigm that a single site mutation is a localised alteration that helps the mechanistic chemist to decipher the chemical mechanism. Thus, much like the T268A mutation, other single-site mutations are not localised alterations but ones that lead to a major change in the complexity of the active site, so as to change the chemoselectivity of reactions.

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