



Cite this: *Phys. Chem. Chem. Phys.*,
2025, 27, 9620

Unusual mechanism of aziridine biosynthesis catalysed by the α KG-dependent non-heme enzyme TqaL[†]

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Aziridines are present in many synthetic pharmaceuticals. The synthesis of the aziridine ring remains challenging due to its highly strained three-membered ring structure. Recently, a non-heme α KG-dependent enzyme, TqaL, has been demonstrated to catalyze the synthesis of aziridines from L-Val. However, the detailed reaction mechanism of the enzyme remains elusive. Herein, we reported, for the first time, the mechanism of oxidative cyclisation for aziridine synthesis catalyzed by TqaL. Following the HAA step, the reaction proceeded *via* a unique concerted process with a single electron transfer from the isopropyl radical to the Fe(III)-OH motif, which was coupled with the electrophilic attack of the primary amine substrate on the tertiary isopropyl radical and simultaneous proton transfer from the substrate amine to the hydroxyl group of the Fe(III)-OH to give the aziridine. This research would provide a valuable structural basis for tailoring the non-heme α KG-dependent enzyme for the biosynthesis of highly active aziridine derivatives as pharmaceuticals.

Received 25th September 2024,
Accepted 1st April 2025

DOI: 10.1039/d4cp03708a

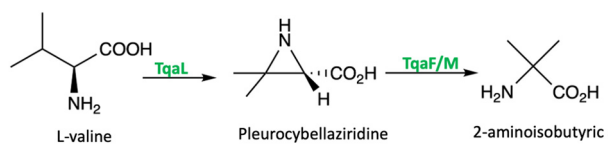
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1 Introduction

Asymmetric synthesis of three-membered nitrogen-containing aziridines is important for the pharmaceutical industry because it is a key building block for many therapeutic drugs with antimicrobial (azicemicins A and B¹), antitumor and anti-malarial activities. Notable examples include mitomycin C, a chemotherapy drug,² and miraziridine A, a potent inhibitor of cathepsin B protease, which is overexpressed in various cancers.³ The synthesis of aziridines represents a challenge for organic chemists due to the high ring strain of the three-membered ring. The biosynthesis pathway of an aziridine natural product catalyzed by engineered heme P450 enzymes^{4,5} and non-heme iron alpha-ketoglutarate (α KG)-dependent enzymes⁶ *via* bimolecular aziridination through a nitrene intermediate has been reported. The bimolecular aziridination catalyzed by iron-containing enzymes has been investigated by theoretical computations.^{7,8} However, to the best of our knowledge, the reaction mechanism for aziridine synthesis *via* unimolecular cyclisation catalyzed by α KG-dependent enzymes has never been reported, which has limited the ability

to tailor the enzyme or discover other novel non-heme enzymes from nature for the synthesis of highly active aziridines.

Recently, the biosynthesis of 2-aminoisobutyric acid by the oxidation of L-Val *via* an aziridine-containing intermediate, pleurocybellaziridine, was reported (Scheme 1). The intramolecular cyclisation is catalysed by the non-heme iron α KG-dependent TqaL enzyme.⁹ Traditionally, reactions catalysed by Fe(II)/ α KG-dependent enzymes proceed through hydroxylation pathways,^{10,11} however, TqaL is a unique non-heme enzyme that catalyzes aziridination rather than hydroxylation. Mutagenesis studies suggested that Phe275 is critical for aziridination over hydroxylation, and mutation of Phe275 to Leu or Ala results in hydroxylated L-Valine as the major product.¹² It was suggested that the aziridination is initiated by the abstraction of a hydrogen atom from the C β of L-Val to the Fe(IV)=O species.^{9,13} Transient kinetics, isotope experiments, combined with product analysis and mechanism probes, indicated that



Scheme 1. The biosynthesis of L-valine to 2-2-aminoisobutyric acid by the action of three enzymes (TqaL, TqaF and TqaM) as reported by Bunno *et al.*⁹ The focus of this study is the first step of the entire process: the oxidation of L-Val to pleurocybellaziridine by the non-heme iron α KG-dependent enzyme TqaL.

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[†] Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4cp03708a>



intramolecular C–N bond formation to install aziridine in L-Val occurs *via* a carbocation intermediate and excluded the possibility of obtaining the final aziridine product *via* a hydroxylated intermediate.¹¹ These experimental hypotheses and evidence enable a detailed understanding of the reaction mechanism of the TqaL enzyme through computational modelling studies at the atomic level. However, in the reported crystal structure of TqaL (PDB: 7EEH), the Fe(IV)=O motif, which is the main oxidant in many Fe(II)/ α KG-dependent enzymes, is absent from the catalytic site, as is the L-Val substrate. The lack of information on how L-Val is positioned in relation to the Fe(IV)=O oxidant has limited our understanding of the aziridination reaction catalysed by TqaL.

Therefore, we employed molecular docking, molecular dynamic simulations (MD) and quantum mechanics and molecular mechanics (QM/MM) calculations to elucidate the configuration of Fe(IV)=O motif and substrate orientation in the active site of the TqaL, and studied the reaction mechanism of aziridine synthesis. Our results show that the C–N bond formation reaction proceeds with the initial hydrogen abstraction from the activated C β of L-valine. Interestingly, following the C–H activation, unlike the mechanism of common α KG-dependent non-heme enzymes that catalyze the hydroxylation *via* the well-known OH rebound step, the reaction catalyzed by TqaL proceeds *via* nucleophilic attack of amine by the electro-negative nitrogen and concerted proton transfer from amine to the Fe(III)–OH, giving the cyclized aziridine product with the release of a water molecule. The research provides structural insights to tailor TqaL for aziridine biosynthesis. The sequence relevance and substrate similarity with hydroxylase (both enzymes can catalyze L-leucine) would enable repurposing of the enzymes or expanding the α KG-dependent non-heme enzyme reservoir in nature.

2 Methods

2.1 Structure preparation

The 3D structure of the enzyme was taken from α KG-dependent dioxygenase TqaL from *Neurospora crassa* (PDB code: 7EEH).⁹ Chain A of the TqaL was used for the computational study. The missing loops of the structure (P97–V116, P172–L180 and G283–N291) were built using the loop modelling protocol (DOPE-HR method) with Modeller software¹⁴ implemented in UCSF Chimera 1.15.¹⁵ The position of α KG was obtained from the α KG analogue *N*-oxalylglycine in the X-ray structure of the homologous enzyme prolyl hydroxylase (PDB code: 6F0W),¹⁶ and then the nitrogen atom in *N*-oxalylglycine was changed to the methylene carbon to give α KG. The water molecule in the vicinity of active site was replaced with superoxo anion radical or oxo complex.

The protonation states of the titratable residues were assigned at pH = 7 using the H++ server.¹⁷ His287 was doubly protonated with an overall charge of +1. The rest of the histidine residues carried an overall neutral charge. His190 and His237 were protonated at the epsilon nitrogen while His148,

His184, His213, His239, and His253 were protonated at the delta nitrogen position. The side chain of all the aspartic and glutamic acids were deprotonated, while all the arginine and lysine residues were protonated. The parameters for L-Val were obtained from the general Amber force field (GAFF). The partial charges were computed using the RESP (restrained electrostatic potential) method by performing quantum mechanical (QM) calculations at the HF/6-31G* level of theory using Gaussian16.

2.2 Molecular docking of L-Val

Flexible docking was performed to dock L-Val in the crystal structures of TqaL using AutoDock 4.2. The structure of the superoxo anion radical was built so that the non-heme iron binds to α KG in a bidentate fashion and to the superoxo anion radical. During the reaction, the superoxo anion radical is decarboxylated to give the Fe(IV)=O species. To understand the location of substrate binding in the active site of TqaL during the reaction, the substrate was docked in both the Fe(III)-superoxide anion radical complex and the Fe(IV)=O complex structures. The grid box was centred at one of the oxygen atoms of the superoxo complex (X, Y, Z: -29.58 Å × 79.286 Å × -8.907 Å) for the Fe(III)-superoxide anion radical complex structure, and the grid box was centred on the oxygen atom of the Fe(IV)=O motif (X, Y, Z: -30.329 Å × 81.691 Å × -7.835 Å) for the Fe(IV)=O complex. In both docking studies, the Lamarckian genetic algorithm (LGA) and the standard free energy scoring function were implemented and a total of 300 LGA runs were carried out for each ligand–protein complex.

2.3 MD simulations

To understand the substrate binding and conformational dynamics of L-Val in the active site of TqaL, MD simulations were performed using the compute unified device architecture (CUDA) version of particle-mesh Ewald molecular dynamics (PMEMD) in Amber20¹⁸ on graphics processing units (GPUs).¹⁹ The parameters for the Fe(IV)=O complex were obtained from our recent research on α KG-dependent thebaine demethylases.¹¹ The Amber ff19SB force field parameters were used for the protein modelling. The protein–substrate complex was then added to the TIP3P truncated octahedral water box, kept at a distance of 15 Å away from the water box edges. Four Na⁺ ions were added to neutralise the protein ligand complex and periodic boundary conditions were used in each simulation run. Particle mesh Ewald was used to compute the long-range electrostatic interactions with a cut-off value of 12 Å and the SHAKE algorithm was used to constrain the bonds involving hydrogen.

The entire system was subjected to two rounds of energy minimization using the steepest descent and conjugate gradient method. In the first energy minimization, the solute molecules were restrained using a potential of 5 kcal mol⁻¹ Å² and the solvent and ions were subjected to 2000-step minimization. In the second round of energy minimization, the entire system was subjected to 5000-step minimization using the steepest descent and conjugate gradient method. The entire system was then heated from 0 to 298.15 K for 50 ps at constant volume using a Langevin thermostat with a collision frequency of 1 ps⁻¹.

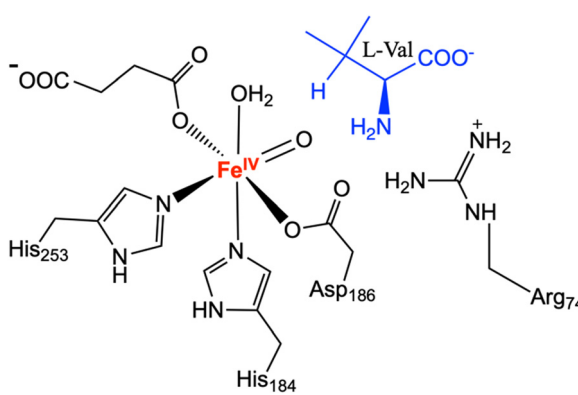


using a canonical ensemble. During the heating process, the non-hydrogen atoms of the solute molecules were restrained using a harmonic potential of 5 kcal mol⁻¹ Å². Then, another 2000-step energy minimization was conducted using the steepest descent and conjugate gradient methods. The equilibration was run at 298.15 K for 50 ps using a weak restraint of 0.1 kcal mol⁻¹ Å² on all the solute atoms in an NPT ensemble and a pressure of 1 bar was maintained using a Berendsen barostat. The 300-ns productive MD simulations were run in the NPT ensemble with a time step of 2 fs. The data was saved every 50 ps and the analysis of MD trajectories were conducted using CPPTRAJ.²⁰

2.4 QM/MM calculations

The reaction mechanism of the aziridine synthesis was studied by performing QM/MM calculations on the most populated snapshots obtained from the MD simulations using ChemShell 3.7.^{21,22} Prior to QM/MM calculations, 2500-step energy minimisation was performed for the selected MD snapshots using the steepest descend and conjugate gradient algorithms. The Fe(IV)=O complex structure was used as the starting point of the reaction for all the QM/MM calculations. The QM region consisted of the side chain of His184, Asp186 and His253 truncated at the C β position, succinate, L-Val and the Fe(IV)=O motif (Scheme 2). Since MD simulations revealed the important role of Arg74 in substrate binding, the effect of including Arg74 in the QM region is considered by additional QM/MM calculations where Arg74 was also added to the QM region.

To perform the QM/MM geometry optimisation, the protein system was split into the relaxed or frozen region. The relaxed region which constitutes protein residues and solvent molecules within 8 Å of iron were allowed to move freely during the geometry optimisation. The rest of the frozen region was kept fixed. The protein residues in the QM region were cut at the C β position and were staturated using hydrogen link atoms. An electronic embedding scheme was used in all the QM/MM calculations to consider the effect of the electrostatic charges of the protein atoms in the MM region on the polarization of the electron density of the atoms in the QM region.



Scheme 2 Quantum mechanics (QM) region in the quantum mechanics/molecular mechanics (QM/MM) calculations.

ORCA 4.2.0²³ was used to run all the DFT calculations with U-B3LYP as the functional of choice.²⁴ The def2-TZVP def2/J auxiliary basis set was used for the iron, while the rest of the atoms were treated with def2-SVP. The D3 dispersion correction and Becke-Johnson (BJ) damping were used in all the calculations. Keywords TightSCF, SlowConv, Grid4, and GridX4 were also used to ensure convergence. For the MM part of the calculations, DL_POLY²⁵ was used with a FF19SB force field.²⁶

To evaluate the ground state of the system, QM/MM geometry optimization was performed for four spin states: singlet ($S = 0$), triplet ($S = 1$), quintet ($S = 2$), and septet ($S = 3$). The QM/MM relaxed potential energy scan (PES) was run for the hydrogen atom abstraction (HAA) step by decreasing the distance (0.1 Å) between the oxygen atom of Fe(IV)=O and the hydrogen atom to be abstracted from L-Val. Starting from the iron oxo oxidant, the reaction to form hydroxylated product *via* the rebound and the possible mechanisms to form the aziridine product include step-wise and concerted intramolecular cyclisation.

The rebound step in the hydroxylation mechanism was calculated by decreasing the distance between the hydroxyl group on the iron centre and the carbon radical on the L-Val with a decrement of 0.1 Å. For the aziridination mechanism, intramolecular attack of the amine on to the C β carbon was modelled by decreasing the distance between the nitrogen and the C β carbon of L-Val with a decrement of 0.1 Å. The transition state associated with each step was fully optimised using the dimer keyword and were then validated by the presence of one unique imaginary frequency. The final energies were also computed at the UB3LYP-D3BJ/def2-TZVPP level.

3 Results and discussion

Most of the reactions catalyzed by non-heme α KG-dependent enzymes (*e.g.*, hydroxylation, demethylation, and epoxidation reactions) follows the common mechanism to first generate an iron oxo oxidant for the subsequent reactions (Fig. 1).^{10,11,27–30,31} The triplet molecular oxygen is coupled to the high spin ferrous iron in the active site of non-heme α KG-dependent enzymes, giving a Fe(III)-superoxide complex (1). The superoxol species is then converted to high-valent Fe(IV)=O species (2) through oxidative decarboxylation of α KG. The Fe(IV)=O motif would then flip to a favourable inline conformation (3) to initiate the oxidation reactions.^{9,32,33}

TqaL is a unique member of non-heme iron α KG-dependent enzymes, as it is the only known enzyme in the family that catalyses the aziridination reaction. In the presence of L-Val, the Fe(IV)=O motif (4) abstracts a hydrogen atom from the C β atom of the substrate to form an isopropyl radical and Fe(III)-OH (5) (Fig. 1).^{12,13} However, unlike the other α -KG dependant non-heme enzymes that produce the hydroxylated product, TqaL catalyzes the aziridination of L-Val to give aziridine.

The formation of aziridine by intramolecular cyclisation may occur from the Fe(III)-OH intermediate (5) *via* three possible pathways (Fig. 1). One possibility is that the reaction proceeds



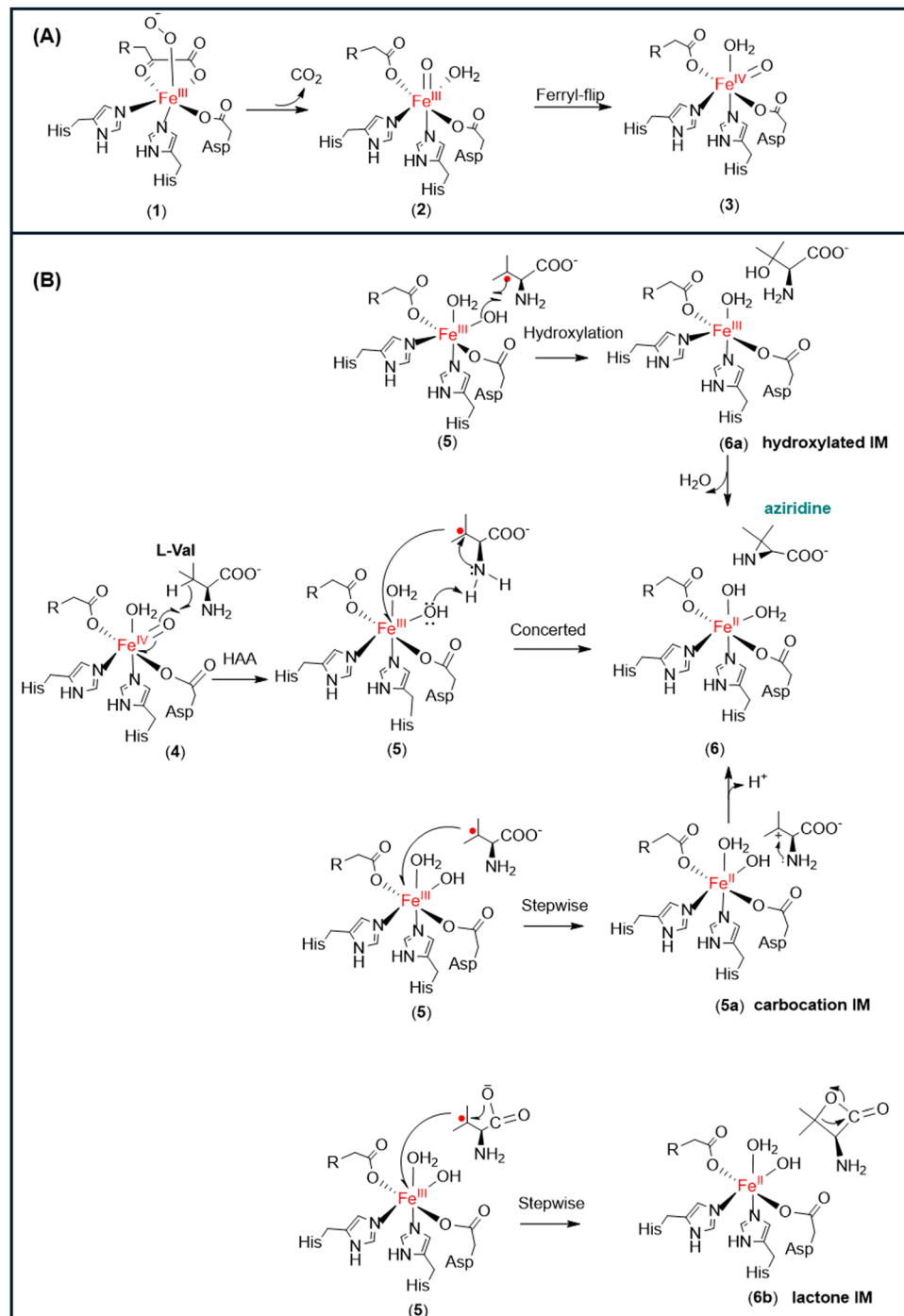


Fig. 1 (A) Formation of the iron oxo oxidant (3) in non-heme α KG-dependent enzymes ($\text{R} = -(\text{CH}_2)_2-\text{COO}^-$). (B) Possible reaction pathways for the formation of the three-membered aziridine from the $\text{Fe}(\text{III})-\text{OH}$ species.

with the rebound of the hydroxyl group to give hydroxylated L-Val (6a), which then cyclises to give the final product aziridine. Another possibility is the transfer of the single electron from the isopropyl radical to the $\text{Fe}(\text{III})$ species to generate a tertiary carbocation intermediate (5a), which is then attacked by the NH_2 group of L-Val to form the aziridine product (6). Alternatively, the single electron transfer and the intramolecular cyclisation may occur simultaneously, which is accompanied by hydrogen atom

abstraction (HAA) from the NH_2 group of L-Val by the $\text{Fe}(\text{III})-\text{OH}$ species to give the final product (6).

3.1 L-Val binding in the active site

The $\text{Fe}(\text{IV})=\text{O}$ species is the main oxidant in non-heme α KG-dependent enzymes,^{10,27,28,30,34-36} therefore, the orientation of $\text{Fe}(\text{IV})=\text{O}$ relative to the substrate in the active site is critical for the enzyme's catalytic capacity. The crystal structure of TqAL

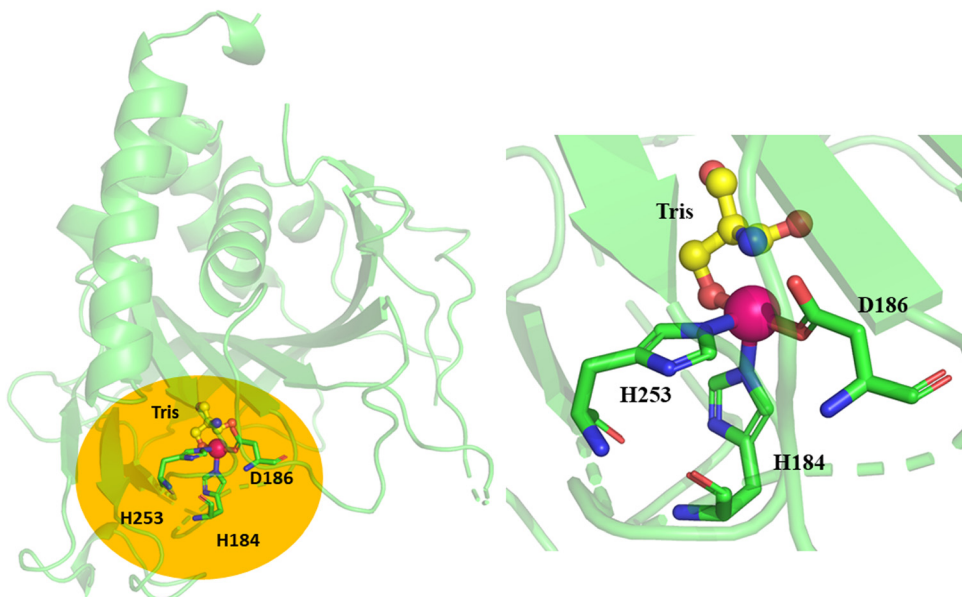


Fig. 2 X-ray structure of chain A of Tqal (PDB code: 7EEH).⁹ The active site of Tqal consists of the mononuclear non-heme iron, shown as a pink sphere, coordinated to 2His/Asp protein residues shown in stick representation. The enlarged catalytic site is shown in the right panel. The Tris buffer is shown in ball-and-stick representations. The missing loops are shown in dotted representation.

(PDB: 7EEH) was used in this study, where the missing loops were modelled using Modeller and then the entire crystal structure was relaxed by 300-ns MD simulations. The structure shows that the mononuclear non-heme iron is coordinated with a triad composed of His184, His253 and Asp186 (Fig. 2). The modelled AlphaFold2 structure is largely same as the crystal structure (Fig. S1, ESI[†]). Hence all the subsequent calculations and analysis are based on the MD refined crystal structure.

The location of L-Val substrate in the active site of Tqal was determined by docking the substrate in the Fe(III)-superoxide and Fe(IV)=O complexes, which correspond to the pre- and post-decarboxylation states, respectively (Fig. 3). Molecular docking studies revealed that Arg74 stabilizes the carboxylate group of the L-Val substrate in the iron oxo complex, and the substrate

binds *trans* to His253, indicating a ferryl flip during the oxidative decarboxylation process (Fig. 3B, see Table S1 (ESI[†]) for other docked poses energies). As a result, the Fe(IV)=O motif rotates and aligns itself toward the C β carbon of the L-Val substrate (the distance between the C β hydrogen of L-Val and the iron oxo oxygen is 2.6 Å), facilitating the HAA to initiate the aziridination reaction. This type of ferryl flip is in agreement with the previous report on other α KG dependent non-heme enzymes.^{27,34,37}

To validate the docking pose and study the stabilization of L-Val in the active site, MD simulations of the Fe(IV)=O state in complex with L-Val (Fig. S2 for the RMSD trajectory, ESI[†]) were performed. The octahedral active site geometry of the active site was maintained throughout the simulations and is consistent with previous literature.^{34,38} It has been established in

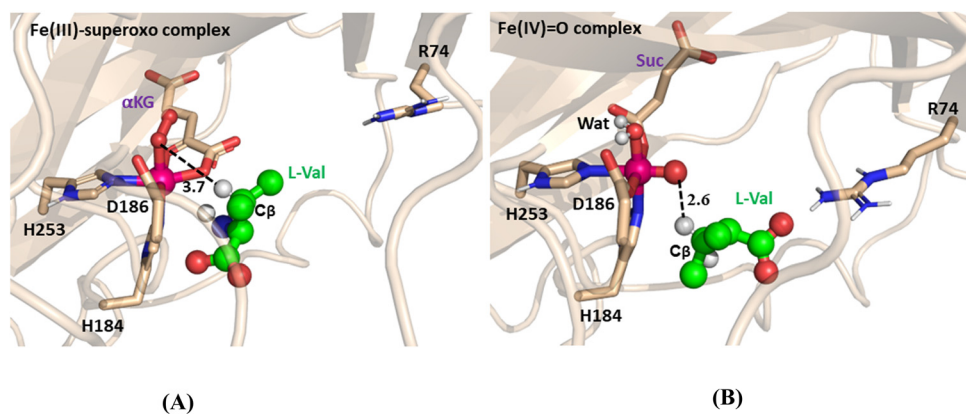


Fig. 3 Molecular docking of the L-Val substrate in the active site of the Tqal enzyme. The docking was performed in both the (A) pre-decarboxylation state (i.e. Fe(III)-superoxide complex) and (B) post-decarboxylation state (i.e. Fe(IV)=O complex) to examine the substrate binding and the configuration of the Fe(IV)=O species.



non-heme and heme enzyme chemistry that the shorter the distance of the hydrogen atom to the oxo group of $\text{Fe}(\text{iv})=\text{O}$ motif, the better it is for catalysis.^{39–41} Recent experimental evidence suggests that aziridination is initiated by the abstraction of hydrogen atoms from the $\text{C}\beta$ of $\text{l}\text{-Val}$ by the $\text{Fe}(\text{iv})=\text{O}$ species.¹³ Therefore, sampling on the MD trajectory was performed based on the distance of the $\text{C}\beta$ to the oxygen atom of the $\text{Fe}(\text{iv})=\text{O}$ complex. Cluster analysis was conducted for the MD simulations and two dominant conformations were selected for further analysis based upon the distance between $\text{Fe}(\text{iv})=\text{O}$ and the $\text{C}\beta$ -hydrogen of $\text{l}\text{-Val}$ (Fig. 4). In both conformations, the carboxylate group of $\text{l}\text{-Val}$ makes electrostatic interactions with the side chain of Arg74 (Fig. S3, ESI†). Our MD simulations show that the isopropyl group of the $\text{l}\text{-Val}$ is stabilized by the hydrophobic residues Phe164 , Ile259 and Phe275 , which is in accordance with the site-directed mutagenesis study, where Phe275 was suggested to be critical for the aziridination reaction and its mutations to Leu or Ala resulted in a hydroxylated major product.¹² MD simulations also showed that the amine group of $\text{l}\text{-Val}$ is well hydrated and forms water-mediated hydrogen bonds with Glu181 . The $\text{C}\beta$ hydrogen atom of the substrate is 2.6 Å from the iron oxo complex, which is an ideal distance to initiate the aziridination reaction by HAA, and agrees with the previous mechanism hypothesis that aziridination is initiated by hydrogen atom abstraction from $\text{C}\beta$ of $\text{V}\text{-Val}$.¹³ However, in the other dominant conformation, the amine group of the substrate moves slightly outwards such that the water-mediated hydrogen bonds with Glu181 breaks (Fig. 4). This results in an increased distance between the $\text{C}\beta$ hydrogen atom and the iron oxo atom (2.6 Å vs 3.0 Å).

To study the mechanism of $\text{l}\text{-Val}$ aziridination, QM/MM calculations were performed for both conformations using the unrestricted B3LYP (U-B3LYP) functional method, which is widely adopted for studying non-heme iron enzymes. A recent study by the Siegbahn group indicates the reliability of B3LYP in capturing the essential electronic properties of metalloenzymes.²⁴ Employing the def2-TZVP basis set for iron ensures a high level of

accuracy in describing the metal center's electronic environment. For the surrounding lighter atoms, the def2-SVP basis set offers a computationally efficient, yet reliable representation. This combination has been extensively utilized in theoretical studies in exploring the mechanisms of non-heme iron enzymes because it provides a balance between computational cost and accuracy. Incorporating D3 corrections effectively improves the description of intermolecular forces in biological systems; hence, it would give more reliable mechanistic insights. The computational settings used in this study are consistent with methodologies employed in recent QM/MM studies of non-heme enzymes.^{34,42,43}

3.2 Aziridination mechanism disclosed by QM/MM calculations

The ground state of the reactant complex was determined by optimising the representative starting structures (Fig. 4) obtained from MD simulations. Four possible spin states of non-heme iron were considered: low-spin open-shell singlet, intermediate spin triplet, quintet, and high-spin septet. Our results show that the spin state order was quintet ($S = 2$) < triplet ($S = 1$) < septet ($S = 3$) < singlet ($S = 0$) (Table S2, ESI†). The quintet is the ground state and is more stable than the triplet spin state by 8.01 kcal mol^{−1}. The singlet and septet spin states are 15–17 kcal mol^{−1} higher in energy compared to the quintet spin state. This is in agreement with the reports for other 2-OG dependent non-heme enzymes for which the quintet is the ground state for the $\text{Fe}(\text{iv})=\text{O}$ complex.^{34,38,44–46} In the electronic configuration of the reactant complex at the quintet state, the spin density of the $\text{Fe}(\text{iv})=\text{O}$ unit is 3.71 and can be described as $\pi^*xy^\dagger \pi^*xz^\dagger \pi^*yz^\dagger \sigma^*x^2-y^2 \sigma^*z^2$. σ^*z^2 is the virtual orbital, and π^*xy , π^*xz , π^*yz and $\sigma^*x^2-y^2$ orbitals are singly occupied with four alpha electrons (Fig. 5A). The electronic structure of $\text{Fe}(\text{iv})=\text{O}$ at the quintet state is consistent with previous theoretical studies on other non-heme αKG -dependent enzymes.^{34,37,47}

3.3 Hydrogen atom abstraction by $\text{Fe}(\text{iv})=\text{O}$

The HAA by the $\text{Fe}(\text{iv})=\text{O}$ unit is well established in various reactions catalysed by non-heme αKG -dependent enzymes such

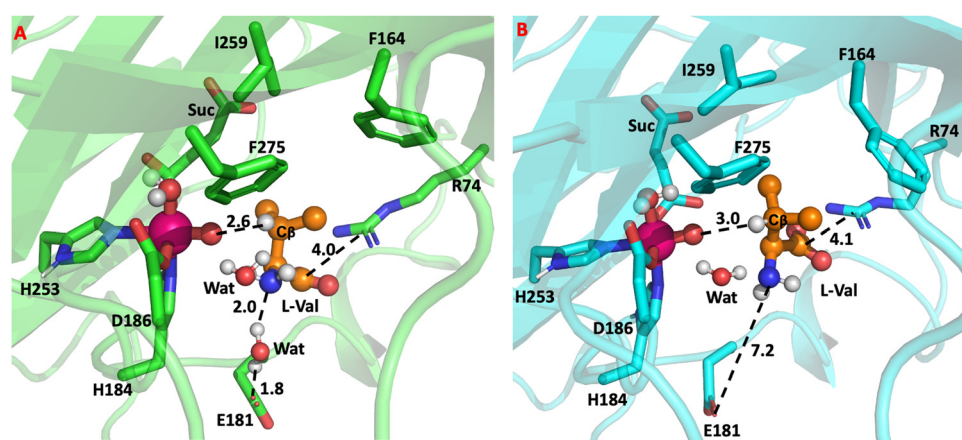


Fig. 4 The most populated conformations (A) and (B) obtained from the MD simulations of $\text{Fe}(\text{iv})=\text{O}$ in complex with $\text{l}\text{-Val}$. The clustering was based on the distance between $\text{Fe}(\text{iv})=\text{O}$ and the $\text{C}\beta$ of $\text{l}\text{-Val}$.



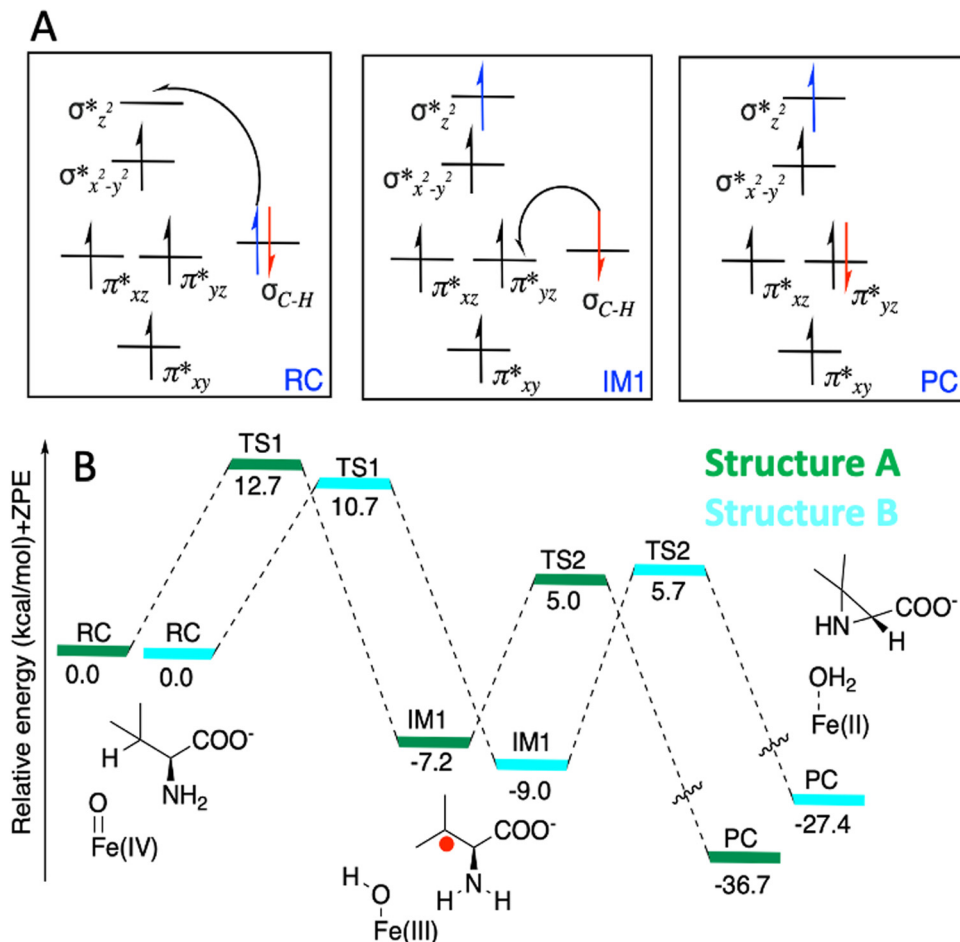


Fig. 5 (A) The electronic configuration obtained during the aziridination reaction catalysed by the TqAL enzyme at the quintet state ($S = 2$). (B) QM/MM reaction profile obtained for aziridine formation via C–N bond formation by intramolecular cyclisation catalysed by the TqAL enzyme at the quintet state ($S = 2$) for both dominant conformations obtained from MD simulations.

as hydroxylation,^{10,28,37,48} epoxidation⁴⁹ or demethylation,^{34,38} however, the mechanism of formation of the cyclised product aziridine initiated through HAA by Fe(IV)=O is unreported. In the reactant complexes for structures A (colour coded green) and B (colour coded cyan) (RC) at the ground state (quintet state), the C β hydrogen of the isopropyl group of L-Val is 2.86 Å and 2.40 Å away from the oxygen atom of the Fe(IV)=O unit, respectively (Fig. 5). The HAA step was also calculated at the triplet ($S = 1$) spin potential energy surface for structure A; however, the high potential energy barrier of 37.2 kcal mol⁻¹ indicates that it is not feasible for the HAA reaction to occur at the triplet surface (Fig. S4, ESI†). Therefore, all the subsequent analysis are for the quintet spin state surface for both A and B structures.

The HAA for structure A and B went through an activation free energy barrier of 12.7 and 10.7 kcal mol⁻¹, respectively and results in the formation of exothermic intermediate IM1 at -7.2 and -9.0 kcal mol⁻¹, respectively (Fig. 5B). In the TS structures obtained from the two starting structures (A and B), the distances between the oxygen atom and the C β hydrogen are 1.36 Å and the distances between the β carbon and the C β hydrogen are 1.25 Å (Fig. 6). The spin density of the Fe(IV)=O

unit increased from 3.71 in RC to 4.01 in TS1 and the spin density of C β decreased from 0 in RC to -0.33 in TS1 (Fig. 6). The change in the spin population and orbital analysis indicates the transfer of the alpha electron from C β of L-Val to the Fe(IV)=O unit and development of the radical character on C β . The electronic configuration of IM1, where the Fe(III)–OH motif has the spin density of 4.55, can be described as $\pi^{*}xy\uparrow\pi^{*}xz\uparrow\pi^{*}yz\uparrow\sigma^{*}x^2-y^2\uparrow\sigma^{*}z^2\uparrow\sigma_{C-H}\downarrow$, and the spin density of -0.96 for C β of L-Val indicates the radical of beta spin on the C β (Fig. 5A). The electron transfer to the Fe(IV)=O unit can either follow a σ or π channel depending on the angle of the C β hydrogen atom facing the Fe(IV)=O motif in the active site. The transfer of an alpha electron to the d_{z^2} orbital indicates the σ pathway and the transfer of beta electron to the $d\pi^*$ orbital known as the π channel. Here, it was observed that the alpha electron was transferred to the $\sigma^{*}z^2$ orbital, indicating that HAA follows a σ pathway. This is in accordance with the HAA mechanism of other non-heme enzymes reported in previous work.⁴⁷

For most non-heme α KG-dependent enzymes, the hydrogen atom is transferred to the iron-oxo motif, followed by the transfer of the hydroxyl group to the substrate radical to give

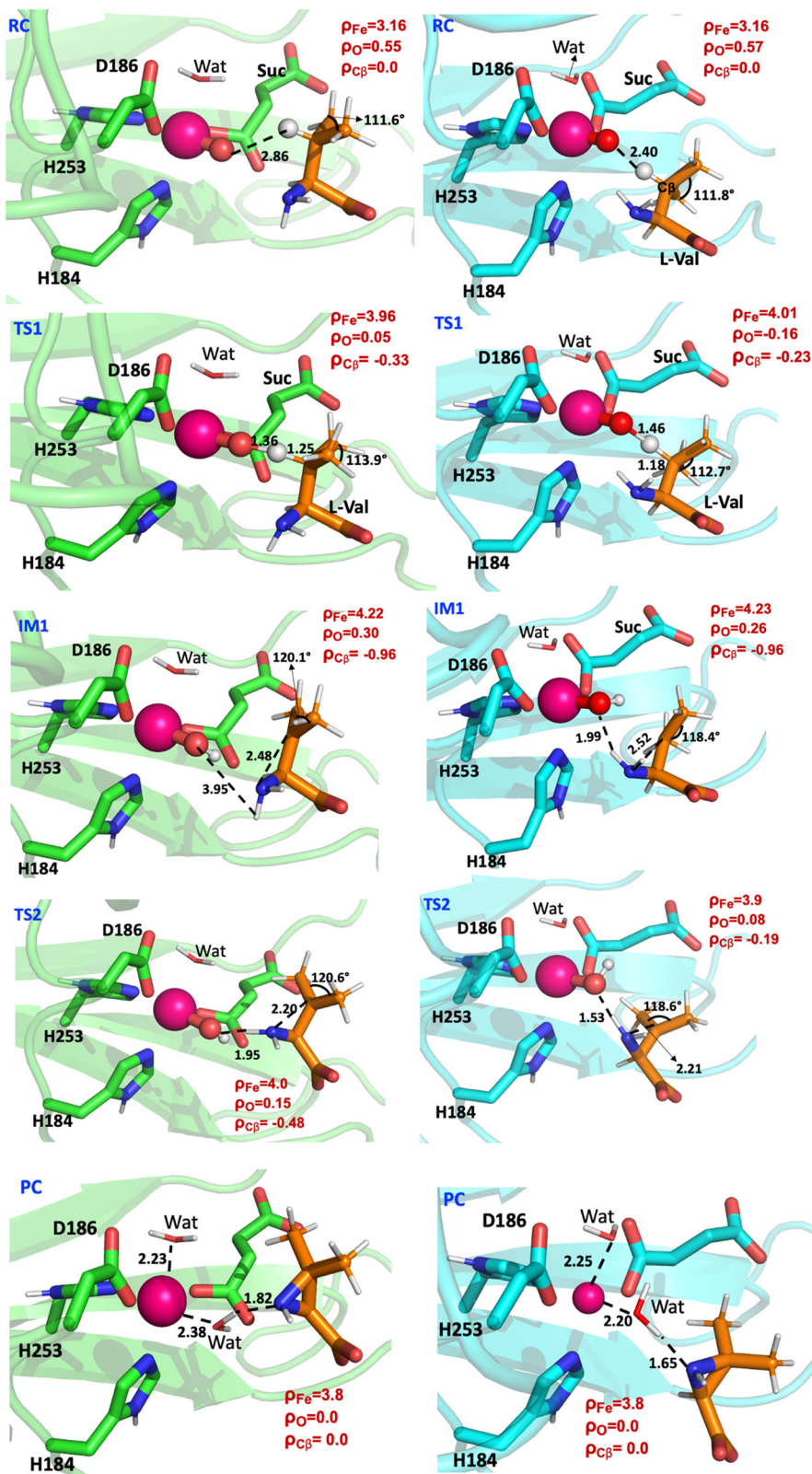


Fig. 6 Stationary and saddle points obtained during the QM/MM calculations of the reaction pathways for the formation of aziridine catalyzed by the TqAL enzyme at the quintet state ($S = 2$) for structure (A) (green) and structure (B) (cyan). The distances are labelled in black text (in Å), and the spin densities (ρ) for each complex are shown in red.



a hydroxylated product *via* a rebound mechanism. However, recent isotope and quantitative production analysis excluded the possibility that the formation of the final aziridine product goes through the hydroxylated L-Val intermediate.¹³ Nevertheless, we still studied the possibility of forming the aziridine product by the rebound mechanism and found that the rebound step goes through the activation energy barrier of 9.6 kcal mol⁻¹, giving hydroxylated L-Val exothermically (Fig. S5, ESI[†]). To obtain the desired product aziridine, the reaction would proceed by transferring the proton from NH₂ to the hydroxyl group of L-Val; however, this step is prohibited with a high potential energy barrier (Fig. S6, ESI[†]). Thus, following the hydrogen abstraction, there must be an alternative pathway, other than the OH rebound mechanism, that the Fe(III)-OH species must follow for the intramolecular cyclisation of L-Val to produce aziridine. Therefore, we investigated other possible reaction mechanisms (*via* a carbocation or lactone intermediate) to form aziridine from L-Val (Fig. 1), as shown in the below section.

3.4 Intramolecular cyclisation *via* a concerted mechanism

It was previously hypothesized that the formation of aziridine goes through a carbocation intermediate rather than the lactone intermediate.¹³ The lactone-based pathway was tested for structure A and it gave the barrier of HAA in excess of 19 kcal mol⁻¹ (Fig. S6, ESI[†]); therefore, it was discarded for future calculations. In the IM1 for structure B (green colour) (Fig. 6), the hydrogen atom of the NH₂ is only 1.99 Å from the hydroxyl group of Fe(III)-OH, so the HAA from the NH₂ group of IM1 by Fe(III)-OH was also studied. However, the potential energy scan shows a high barrier of 24.3 kcal mol⁻¹ (Fig. S7, ESI[†]), indicating that it is unlikely for the reaction to proceed with HAA from the amine group. We also considered the possible stepwise mechanism that includes a single electron transfer from the isopropyl radical to form the potential carbocation 5a (Fig. 2B), which would then undergo intramolecular cyclisation to give the aziridine product. However, all the optimisation attempts to obtain 5a reverted back to 5, further underpinning that the reaction proceeds *via* an alternative mechanism.

During the HAA, the angle of the isopropyl group increased from 111.6° in the initial reactant complex (RC) to 120.1° in the IM1 (Fig. 6), which corresponds to a trigonal planar geometry of the isopropyl group and resembles the conformation of a tertiary carbon radical. The trigonal planar geometry of IM1 is crucial, as it allows for IM1 to act as a Lewis acid, with its partially occupied p orbital on the C β to orient itself perpendicular to the plane of the isopropyl group, and therefore enable the intramolecular cyclisation to form aziridine. The NH₂ in IM1 is only 2.48 Å from the C β atom of L-Val, which facilitates nucleophilic attack of the planar C β of L-Val and the intramolecular cyclisation of L-Val. This process is coupled with a single electron transfer of beta spin localised on the C β atom to the central iron atom, going through the transition state TS2 with an activation energy barrier of 12.2 kcal mol⁻¹ (Fig. 5B).

The spin density of Fe(III)-OH decreases from 4.52 in IM1 to 4.15 in the transition state TS2 and that of C β of L-Val increases

from -0.96 to -0.48. In the TS2 structure, the distance of NH₂ to the C β atom of L-Val is 2.20 Å and the hydroxy group of Fe(III)-OH is 1.95 Å away from the proton of NH₂ group. The intramolecular cyclisation occurs *via* a concerted mechanism with nucleophilic attack of NH₂ of the L-Val to the C β atom and the transfer of the proton from NH₂ to the hydroxy group of Fe(III)-OH. During the concerted process, a water molecule leaves to exothermically yield an aziridine product at -36.7 kcal mol⁻¹ (Fig. 5B).

4 Conclusions

TqaL offers an attractive biosynthetic route for the incorporation of three-membered aziridine ring structures into the core of biologically active molecules. It is critical to decide the binding of the substrate in the enzyme for rational engineering to improve the substrate scope to potentiate its use for the pharmaceutical industry. However, the substrate is missing in the crystal structure of the enzyme and the reaction mechanism of aziridine synthesis *via* unimolecular cyclisation catalyzed by a non-heme enzyme has never been reported.

Here, we reported that the binding of L-Val in the active site of TqaL is stabilised by Arg74, Phe164, and Phe275. These residues enable the C β hydrogen of the L-Val substrate to be abstracted by the Fe(IV)=O motif. The resulting Fe(III)-OH species then accepts an electron from the C β radical of the substrate. The single electron transfer is accompanied by the attack of a polar amine to the carbon radical and simultaneous transfer of a proton from the amine group of the substrate to the Fe(III)-OH species to form the final aziridine product. The active conformation for substrate binding and the elucidation of the uni-molecular cyclisation for aziridine synthesis sets the basis for engineering non-heme enzymes to exploit their use in the biosynthesis of aziridine-derived active therapeutics.

Data availability

The data supporting this article have been included as part of the ESI.[†]

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We are grateful for the computing resources from the QUB high performance computing Tier2 resource, funded by EPSRC (EP/T022175). W. S. acknowledges the support of Research England's Expanding Excellence in England (E3) Fund.

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