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Oxygen diffusion pathways in a cofactor-independent dioxygenase†

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Molecular oxygen plays an important role in a wide variety of enzymatic reactions. Through recent research efforts combining computational and experimental methods a new view of O₂ diffusion is emerging, where specific channels guide O₂ to the active site. The focus of this work is DpgC, a cofactor-independent oxygenase. Molecular dynamics simulations, together with mutagenesis experiments and xenon-binding data, reveal that O₂ reaches the active site of this enzyme using three main pathways and four different access points. These pathways connect a series of dynamic hydrophobic pockets, concentrating O₂ at a specific face of the enzyme substrate. Extensive molecular dynamics simulations provide information about which pathways are more frequently used. This data is consistent with the results of kinetic measurements on mutants and is difficult to obtain using computational cavity-location methods. Taken together, our results reveal that although DpgC is rare in its ability of activating O₂ in the absence of cofactors or metals, the way O₂ reaches the active site is similar to that reported for other O₂-using proteins: multiple access channels are available, and the architecture of the pathway network can provide regio- and stereoselectivity. Our results point to the existence of common themes in O₂ access that are conserved among very different types of proteins.

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Introduction

The diffusion of molecular oxygen (O₂) is an essential process in living organisms, where it plays an important role in a large variety of enzymatic reactions. Although it was initially believed that O₂ freely diffused through proteins,¹ recent research efforts uncovered the existence of dynamic gas molecule channels that grant temporal and spatial regulation,^{2–11} and provide insight into the known regio- and stereo-selectivity of some oxygenase-catalyzed reactions.^{4,12}

In most cases, permanent channels are not observed in static protein structures as diffusion occurs through a network of transient pathways that are mostly lined with flexible hydrophobic residues.^{2,4,13,14} In addition, O₂ is not usually observed in protein crystal structures and although xenon can sometimes successfully identify O₂ binding sites,^{5,7,15,16} it is far from an ideal probe.¹⁷ Nonetheless, the problem of O₂ diffusion can be ideally addressed using computational tools. So far, computational studies of O₂ diffusion have mainly relied on implicit ligand sampling,^{4,11} multiple steered molecular dynamics¹⁸ or enhanced sampling techniques^{2,3} to attain adequate sampling. However, in recent years GPU computing produced a large

increase in the simulation timescales that can be routinely accessed,^{19–22} making it possible to obtain the several independent ~100 ns long simulations that are necessary to achieve a realistic description of ligand motion using straightforward molecular dynamics.² In this work, we show how this approach can be applied to study O₂ diffusion in a cofactor-independent oxygenase.

The majority of enzymes that catalyze reactions where oxygen atoms from O₂ are transferred to organic substrates require metals or organic cofactors like flavin to activate O₂ and control the reactivity of the intermediates formed.¹² However, a small number of oxygenases require neither.²³ DpgC, an enzyme that catalyzes a key step in the biosynthesis of a non-proteinogenic amino acid that is incorporated into last-resort antibiotics like vancomycin and teicoplanin, belongs to this rare group.^{24–26} Because of emerging bacterial resistance,²⁷ these antibiotics have been the focus of numerous biosynthetic and synthetic chemistry efforts.^{28–32} Interestingly, DpgC is one of the rare cases where a proposed O₂ binding site was apparent in the crystal structure, in this case located in a solvent accessible hydrophobic cavity, and positioned to attack only a specific face of the substrate (3,5-dihydroxyphenylacetyl-coenzyme A, DPA-CoA),³³ in agreement with labeling experiments.^{26,33}

Here, we investigate O₂ diffusion in DpgC using long, unbiased molecular dynamics simulations that reveal the importance of including all subunits of this hexameric protein to obtain a realistic representation of O₂ diffusion pathways. Site-

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directed mutagenesis, kinetic measurements, and Xe-pressure-crystallography provide information validating the computational results and supporting the existence of multiple O₂ diffusion pathways that converge at the active site face where O₂ was crystallographically observed. These observations point to the existence of common features in O₂ access that are conserved among many different types of proteins.

Results

Molecular dynamics simulations reveal three main oxygen diffusion pathways in DpgC

We investigated the diffusion of O₂ into and out of the active site of DpgC using molecular dynamics simulations. These started from the X-ray structure of the hexamer, with O₂ located in the active site position observed crystallographically (PDB: 2NP9, 2.45 Å resolution). Each of the five independent simulations contained six O₂ molecules and extended for over 0.5 μs of combined time. During these simulations, we observe 55 complete spontaneous diffusion events that bring O₂ from outside the protein into the active site, and 72 escapes. Complete entries are defined as events where O₂ starts completely outside the protein surface and ends in a position less than 6 Å away from the reactive carbon of the substrate, DPA-CoA, with the opposite being true for complete escapes (see Fig. S1† for a scheme of the reaction). On average, each O₂ molecule spends 20% of the trajectory traveling along a complete path. We consider that these simulations provide sufficient sampling, as the main results are in reasonable agreement with an additional set of simulations where the concentration of O₂ was approximately six times higher and the initial position of all the O₂ molecules was outside the protein (see ESI† for more details).

The observed spontaneous diffusion of O₂ takes place through well-defined pathways (Fig. 1). There are three main types of pathways: the U1 pathway, in which O₂ enters the

enzyme through the *upper* part of the active site; the S1 pathway, which connects the active site with a region located at the interface of two of the *subunits*; and the L-type pathways, in which O₂ reaches the active site by diffusing close to the *ligand* DPA-CoA.

Among the three main types of pathways identified, the L1 pathway accounts for the majority of all observed entries and escapes (Fig. 2). In total, O₂ escaped through this pathway 46 times and entered 39 times. To enter the L1 pathway, O₂ diffuses between Lys 428 and Phe 432 (Fig. 3A). The entrance to the L2 pathway is very close to that of the L1 pathway and is located between Lys 428 and the loop formed by residues 318 to 322. We observed other L-type pathway diffusion events with low frequency. In the U1 pathway, O₂ diffuses through a hole lined by hydrophobic residues located at the top of the active site (Fig. 3B). The size of the hole across its smallest dimension (Ala 112 – Val 429) is 6 ± 2 Å. Finally, the S1 pathway connects the active site to a region found at the interface of two of the protein subunits. Although only one complete entry and escape occurred *via* this pathway, O₂ molecules regularly visited several cavities along this pathway without completing an escape or entry during the simulation. It is important to point out that because of the location of the entrance to this pathway, O₂ may need to diffuse up to ~50 Å between DpgC subunits before reaching this point (Fig. 3D). This highlights the importance of simulating the hexamer in order to obtain a realistic representation of O₂ diffusion, as already pointed out by Shadrina *et al.* in simulations of hemoglobin where the enhanced flexibility of isolated subunits increased O₂ migration.² This concept is even more relevant in DpgC where, unlike in hemoglobin, we observe O₂ diffusion between protein subunits.

O₂ concentrates in a dynamic cavity at the active site of DpgC

When O₂ travels along any of these pathways it is usually found in one of several pockets, which are connected through bottleneck regions (Fig. 4A). The lower pocket is connected to the access points for the L1 and L2 pathways, while the upper pocket is connected to the U1 access point. O₂ molecules diffusing along the S1 pathway occupy pockets SI–SIV. When all complete paths are considered, even though the time spent in each pocket can vary in a wide range, the L1 pocket is the one that is populated the most. This pocket is connected to the

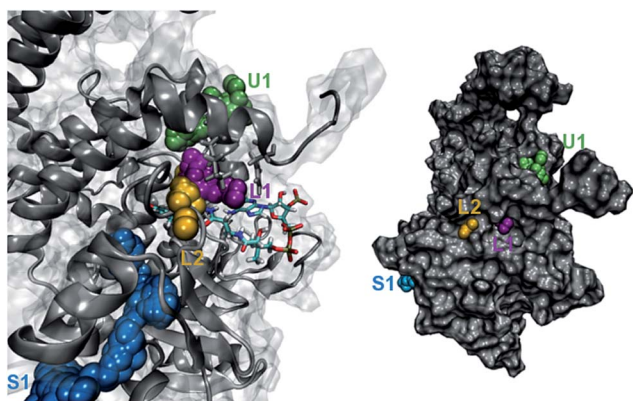


Fig. 1 Main pathways for O₂ diffusion in DpgC. Green: U1 pathway, purple: L1 pathway, yellow: L2 pathway; blue: S1 pathway. Left: trajectory followed inside the protein. Right: entry and escape points shown on the surface of one of the monomers. A video is included in the ESI.†

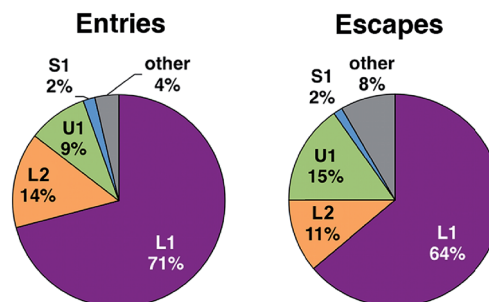


Fig. 2 Distribution of O₂ pathway use for complete diffusional processes.



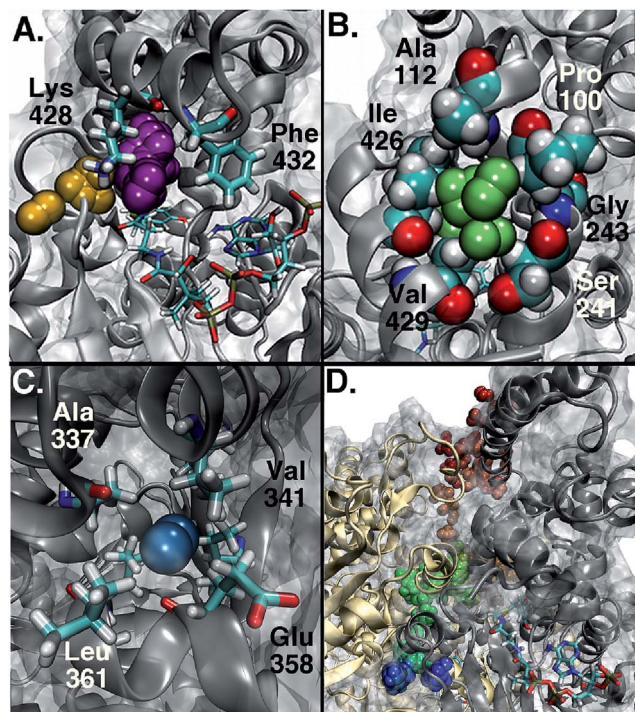


Fig. 3 Entry points to the different types of pathways. (A) In the L1 pathway, O_2 diffuses between Lys 428 and Phe 432. (B) In the U1 pathway, O_2 diffuses through a hole lined by hydrophobic residues shown in space filling representation. (C) Residues forming the entry point to the S1 pathway. (D) The entry point to the S1 pathway is located at the interface between two of the subunits, shown in different colors. In order to reach this point, O_2 needs to diffuse between them. An example of such a trajectory is shown, with the position of O_2 along it progressing from red to blue.

upper pocket, and SI. Only when O_2 is in the lower or SI pockets it can be found at a distance of less than 6 Å from the carbon atom of DPA-CoA where it is expected to react. On average, O_2 molecules spent 3% of a trajectory in this position. As a result, the probability of finding O_2 in a reactive position is 10 to 60-fold higher than if O_2 was uniformly distributed, as would be expected from passive diffusion. This observation is in good agreement with the assigned oxygen binding pocket in the crystal structure. It is also important to point out that O_2 only has access to DPA-CoA from one face, in agreement with labeling experiments that suggest stereochemical hydrogen abstraction, and O_2 attack from the opposite face.^{26,33} In addition, water is relatively excluded from the active site except for the crystallographically observed waters hypothesized to act as a base in the reaction mechanism (Fig. S2†).

The pockets where O_2 is found during complete entries and escapes are dynamic. Analysis by a geometry-based cavity detection algorithm (MDpocket³⁴) reveals that the lower pocket is available roughly 50% of the time in all protein subunits, while the upper pocket is only available 10–40% of the time. SIII is also available 50% of the time in all subunits, while the rest of the pockets along the S1 pathway display more variability between subunits (see Fig. S3† for more details). In addition to these pockets, MDpocket also identified several other cavities,

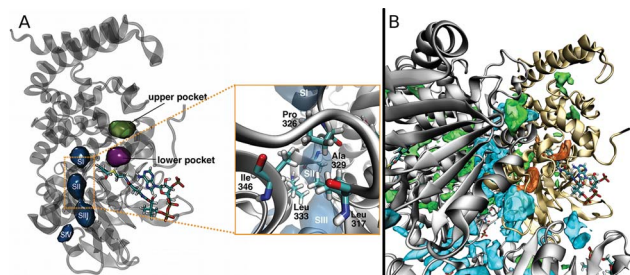


Fig. 4 (A) Pockets where O_2 is found along complete paths, representation generated based on O_2 occupancy. (B) Cavities identified by MDpocket as being available 50% of the time. Orange: pockets where O_2 resides along complete entries and escapes. Green: other pockets located inside a single subunit. Blue: pockets located between subunits. One of the subunits in the hexamer is highlighted in yellow for reference.

shown in Fig. 4B, that are frequently populated by O_2 in our simulations. Many of these are located between protein subunits and play an important role in allowing O_2 access to the entry point of the S1 pathway.

Implicit Ligand Sampling (ILS) post-processing of MD simulations provides further insight into the free energy landscape of the O_2 diffusion pathways. This method evaluates the energy associated with placing an O_2 molecule at any point of a 3-dimensional grid.³⁵ Fig. 5 summarizes the free energy profiles obtained for each pathway, with all values relative to O_2 in the solvent. The lower and SII pockets display the lowest free energy, but all pockets stabilize O_2 with respect to the solvent. The free energy of the lower pocket ($-4.3 \text{ kcal mol}^{-1}$) is comparable to the one observed for the active site of lipooxygenase ($-4.7 \text{ kcal mol}^{-1}$).⁴ It was also possible in all cases to identify hydrophobic surface sinks that concentrate O_2 at the pathway gates, in agreement with other computational studies.^{3,4} For pathways L1, L2, and U1 the barriers for O_2 entry were in the 1.2 to 1.6 kcal mol^{-1} range, also similar in value to those observed in lipooxygenase using the same method.⁴ In agreement with the molecular dynamics results, the lowest barrier of the three was observed for L1, and the highest for U1. In contrast, the S1 pathway displays larger free energy barriers and in particular the one for the transition between SII and SI is $\sim 10 \text{ kcal mol}^{-1}$. This very large barrier is consistent with the low frequency for the S1 pathway in the MD simulations, where we observe the transition between these pockets only twice, but we see O_2 in the rest of the pockets along the S1 pathway more frequently. It is important to note that due to the high barrier observed for this pathway, ILS alone would probably not have identified it as a viable access channel.

F432W and F432Y hinder O_2 diffusion along the L1 pathway

Since the L1 pathway was observed with a high frequency, we expect mutations that alter access to this pathway to have a significant impact on kinetic parameters affected by O_2 diffusion rates. Using site-directed mutagenesis, we targeted F432 and mutated it to a residue with larger steric bulk (F432W), to a smaller amino acid (F432L), and to tyrosine. In the F432Y



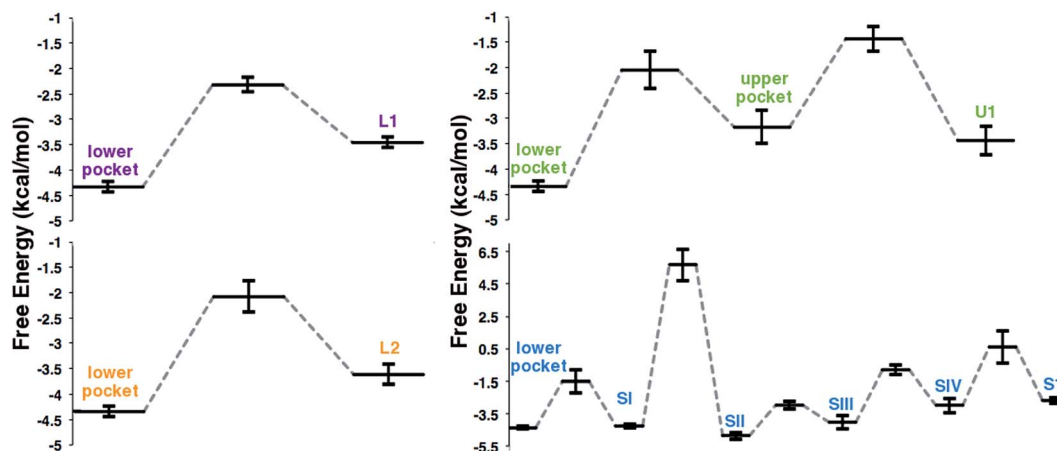


Fig. 5 Free energy profile for O_2 migration along pathways L1, L2, U1, and S1. Free energies are relative to O_2 in the solvent. Error bars correspond to the standard error for the six different subunits.

mutant, the interactions between the phenol and DPA-CoA anchor the mutated residue producing a loss of mobility that hinders access through the L1 pathway. Short molecular dynamics simulations of the mutants reveal that in F432W the percentage of time the L1 pathway gate is open decreases by $\sim 10\%$, and by $\sim 15\%$ in F432Y (Table S8†).

All mutants exhibited a drop in k_{cat}^S and K_M^S in kinetic measurements with varying DPA-CoA concentrations (Table 1). F432L retained less than 20% of the WT's catalytic activity and displayed a $\sim 50\%$ decrease in the DPA-CoA binding constant driven by a reduction in the binding enthalpy (Table S10†). Since the mutation affected substrate binding, F432L cannot be used to evaluate the effect of altering O_2 access to the L1 pathway and was ruled out from further consideration. For both F432W and F432Y k_{cat}^S and K_M^S decrease in the same proportion, as expected for mutations that decrease the rate of O_2 diffusion into the active site (see ESI†). Kinetics experiments with varying concentrations of O_2 confirm this observation, revealing a significant decrease in $k_{cat}^{O_2}/K_M^{O_2}$. Assuming that the mutations did not affect the catalytic activity of DpgC this represents a $\sim 300\%$ increase in $K_M^{O_2}$, consistent with a large decrease in the conductivity of the L1 pathway. In agreement with our mutant simulations, the decrease in $k_{cat}^{O_2}/K_M^{O_2}$ is larger in F432Y due to the larger impact of this mutation on the percentage of time that this pathway is accessible.

Xenon binds along the S1 pathway

According to the results of our molecular dynamics simulations, O_2 diffusion along the S1 pathway takes place with low probability and, as a consequence, we expect mutations that alter this pathway to have a relatively small impact on kinetic parameters affected by O_2 diffusion rates. Using site-directed mutagenesis, we substituted a residue at the S1 gate for a less sterically hindering one expected to increase conductivity through the pathway (L361A), and an amino acid located near the bottleneck region between the SII and SIII pockets for a bulkier one (A329L, Fig. 4A).

The L361A mutant exhibited an increase in both k_{cat}^S and K_M^S in kinetic measurements with varying DPA-CoA concentrations (Table 1) but these did not increase by the same factor, as expected from a mutation that increases the rate of O_2 diffusion without affecting DPA-CoA binding or catalysis (see ESI†). While K_M^S is approximately 2 times larger than that for WT DpgC and the k_{cat}^S is 20% larger, these increases are not statistically significant. In line with these observations, kinetic measurements with varying O_2 concentrations produced results that are indistinguishable from those of the WT protein.

The A329L mutant retained only $\sim 10\%$ of the WT catalytic activity, while its K_M^S is approximately 3 times larger. Such a drastic reduction in k_{cat}^S is unexpected from a conservative, non-

Table 1 Kinetic parameters for WT DpgC and different mutants affecting O_2 access to the L1 pathway. The error estimation corresponds to the 95% confidence interval

Mutant	DPA-CoA kinetics			O_2 kinetics
	K_M^S (μM)	k_{cat}^S (s^{-1})	k_{cat}^S/K_M^S ($\mu M^{-1} s^{-1}$)	$k_{cat}^{O_2}/K_M^{O_2}$ ($\mu M^{-1} s^{-1}$)
WT	2.3 ± 1.8	0.15 ± 0.02	0.07 ± 0.02	2.2 ± 0.1
F432W	1.1 ± 0.4	0.081 ± 0.006	0.07 ± 0.01	0.81 ± 0.05
F432Y	1.9 ± 0.6	0.125 ± 0.009	0.07 ± 0.01	0.52 ± 0.05
F432L	0.5 ± 0.2	0.03 ± 0.02	0.06 ± 0.02	N/A
A329L	6.6 ± 1.9	0.017 ± 0.001	0.0026 ± 0.0004	1.1 ± 0.2
L361A	4.4 ± 1.1	0.18 ± 0.02	0.042 ± 0.005	2.1 ± 0.2



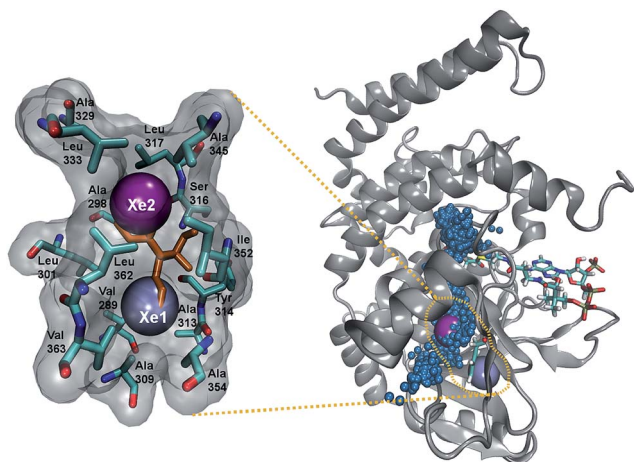


Fig. 6 Left: Residues lining the Xe-binding sites, hydrogen atoms not shown. The two observed conformations of F315 are shown in orange. Right: location of the Xe-binding sites within one of the DpgC subunits. The blue dots represent the positions of O_2 along a trajectory where it escaped using the S1 pathway. F315 and DPA-CoA shown for reference.

active site mutation targeting an infrequently used pathway, especially considering the minimal impact of the L361A mutation. Kinetic measurements with varying O_2 concentrations exhibit a significant decrease in $k_{cat}^{O_2}/K_M^{O_2}$. Assuming that the mutation did not affect the catalytic activity of DpgC, this represents a 100% increase in $K_M^{O_2}$, implying a large decrease in the conductivity of the L1 pathway. However, the large unexpected loss of catalytic activity caused by the mutation suggests that this might not be a justified assumption.

Additional insight into the role of the S1 pathway in O_2 diffusion comes from X-ray diffraction experiments applying Xe as an O_2 probe. Using Xe to study gas migration is a powerful tool because its anomalous signal allows clear identification and its polarizability and van der Waals radius are comparable to those of diatomic molecules like O_2 , CO, and NO.^{5,7} Exposing DpgC crystals to 300–500 psi of Xe revealed two binding sites, none of them corresponding to the crystallographically observed O_2 binding site (Fig. 6 PDB: 4YLH). The minor site (Xe2) is located in the hydrophobic SIII pocket, which according to our MDpocket calculations is the only pocket along the S1 pathway that is available at least 50% of the time. This site is regularly visited by O_2 during complete trajectories along the S1 pathway. In contrast, O_2 does not populate the major Xe binding site, which can only be accessed by displacing F315 from its typical conformation. In our simulations, F315 is part of the boundary of the SIII pocket and prevents O_2 from diffusing closer to the protein surface, adopting the alternative conformation only 9% of the time (Fig. S9†). ILS calculations reveal that the Xe1 and Xe2 sites display a binding free energy of -4.6 and -4.4 kcal mol⁻¹ respectively (see ESI† for more details). These, combined with the high hydrophobicity of the sites, explain why Xe binds to these sites only and not to the active site, which is more polar and solvent-accessible. Similar observations were made for cofactorless urate-oxidase, where

the relatively polar O_2 binding site in the active site does not match the Xe binding sites.¹⁷

Discussion

Pathways and gates for ligand diffusion – simulation vs. experiment

Based on the results of our molecular dynamics simulations, O_2 diffusion in DpgC occurs through three main pathways with four distinct access points. In general, computational studies of small gas molecule diffusion in proteins also find multiple pathways and gates.^{3,4,6,14,36,37} However, these results must be interpreted carefully. Ligand diffusion in myoglobin has been extensively studied, and most experiments agree on diffusion taking place 70–80% of the time through a single access point known as the histidine gate.¹⁴ While there is good agreement between simulations and experiments with respect to identifying ligand-binding cavities and connections between them, simulations usually predict the existence of multiple gates. Ruscio *et al.* used molecular dynamics simulations to identify two main CO diffusion pathways, which connected all Xe-binding sites, with nine total gates.⁶ While the simulations correctly identify the histidine gate as the most frequently used access point, it only accounts for ~30% of all complete entries and escapes. Although the multiple pathways and gates model does not provide a description of small ligand diffusion that fully agrees with experimental results for globins, it seems to be a good representation for other proteins such as copper-containing amine oxidase, where mutations blocking individual pathways have little effect on $k_{cat}/K_M(O_2)$.³⁸

In the case of DpgC, our simulations point to one pathway and gate accounting for a large majority of the complete entries and escapes observed and, in agreement with this observation, mutations that hinder access into L1 reduce $k_{cat}^{O_2}/K_M^{O_2}$. While mutating the gate to the S1 pathway had negligible effect, the conservative A329L mutation had an unexpectedly large effect. This mutation blocks the SIII pocket and, as a result, the minor Xe-binding sites. The SIII pocket is available at least 50% of the time during our simulations and is connected to the other pathways at the active site. Taken together, this data suggests the possibility of the SIII pocket acting as a temporary O_2 storage. A complete assessment of the importance of the different pathways would require mutations along U1 and L2. We evaluated different possible mutations using short molecular dynamics simulations but found that these altered the protein structure affecting more than one pathway at a time (see ESI† for more details).

O_2 pathways guide regio- and stereospecificity

According to our simulations, in DpgC O_2 only has access to DPA-CoA from one face, in agreement with labeling experiments that suggest stereochemical hydrogen abstraction, and O_2 attack from the opposite face.^{26,33} Other enzymes also exert regio and stereoselectivity by guiding O_2 to specific positions within the active site.³⁷ Implicit ligand sampling simulations of 12/15-lipoxygenase show that oxygen concentrates around a



specific carbon atom in a position that promotes O₂ insertion from the opposite face as hydrogen abstraction.⁴ Enhanced-statistics molecular dynamics in two different flavoenzymes revealed funnel-like pathways that guide O₂ to the re-side of the flavin C4a atom.³ Thus, the positions and connections of the O₂ access channels play a key role in determining the identity of the reaction products in metal-dependent, flavin-dependent and cofactor-dependent oxygenases suggesting that this strategy is conserved among many different types of enzymes.

Binding vs. consuming O₂: avoiding enzyme inactivation

O₂ activation takes place in a series of one-electron reactions producing highly reactive radical intermediates that can cause oxidative inactivation. Enzymes evolved a number of strategies to minimize the damage caused by these side-reactions.¹² As mentioned above, O₂ diffusion channels provide regio- and stereochemical control of the reaction products. In addition, most enzymes use metals or organic cofactors such as flavin to control the reactivity of the oxygen intermediates. In particular, many flavoenzymes display very low O₂ binding affinities with kinetics that show no saturation with O₂, effectively consuming O₂ without binding it tightly.³⁷

DpgC is a cofactor-independent dioxygenase displaying negligible oxidative inactivation and reaction side-products and as such can provide interesting insight into how proteins control O₂ reactivity in the absence of metals and organic cofactors. Reactive oxygen species have been implicated in the mechanism of DpgC and other cofactor-independent oxygenases.^{23,26,33,39,40} Sulfur-containing amino acid residues are particularly sensitive to oxidation by reactive oxygen species.⁴¹ Interestingly, in DpgC none of these residues are located along the diffusion pathways found to lead O₂ into the active site (Fig. S5†). In addition, the number of entries and escapes observed in DpgC are ~10 times higher than those reported in molecular dynamics simulations of myoglobin⁶ and flavin-dependent enzymes³ (Table S7†), after accounting for differences in simulation times and O₂ concentrations. However, although DpgC concentrates O₂ in the active site by 10–60 times compared to having O₂ uniformly distributed, this is significantly less than in the case of lipoyxygenase (50–100 times)⁴ and flavin-dependent D-amino acid oxidase (~400 times).¹¹ Taken together, this data suggests that in the absence of metals or cofactors, enzymes such as DpgC may control O₂ reactivity through weak binding that minimizes unproductive O₂ residence times in the active site, compensated with a higher frequency of O₂ entries.

Conclusions

The results of our computational simulations, supported by the mutagenesis experiments and Xe-binding data provide direct evidence for the existence of functional O₂ diffusion pathways in a cofactor-independent oxygenase. Although DpgC is rare in its ability of activating O₂ in the absence of cofactors or metals, the way this gas molecule reaches the active site is similar to what other authors report for flavoenzymes,³ metal-dependent

oxygenases⁴ and even globins:⁶ multiple pathways can be involved, and the architecture of the pathway network can play an important role in controlling regio- and stereoselectivity. These results point to the existence of common themes in O₂ access that are conserved among very different types of proteins.

O₂ diffusion in DpgC takes place through three main pathways with four access points, with the L1 pathway accounting for most of the observed diffusion events in our simulations. These pathways guide O₂ to a specific face of DPA-CoA, in agreement with labeling experiments that point to stereochemical hydrogen abstraction.^{26,33} The pathways connect a series of dynamic hydrophobic pockets. The lower pocket, where O₂ was crystallographically observed, concentrates O₂ and positions it for reaction with DPA-CoA. Our computational results highlight the importance of simulating the complete biological unit whenever possible, as diffusion between protein subunits is important to obtain an accurate description of the S1 pathway. In addition, molecular dynamics provide unbiased O₂ diffusion pathways, as only the protein and solvent guide O₂ motion, granting information about the relative importance of each pathway that is hard to obtain from cavity-detection methods.

Methods

Molecular dynamics simulations

All simulations were performed using the AMBER12 suite.⁴² The ff99SB force field parameters were used for all residues⁴³ except DPA-CoA. Parameters for this molecule were generated combining published parameters for adenosine triphosphate⁴⁴ and HF/6-31G* Gaussian09⁴⁵ calculations. The five non-modeled C-terminal residues in the crystal structure of DpgC (PDB: 2NP9, 2.45 Å resolution) were added manually and the non-hydrolyzable substrate-analogue was substituted with substrate DPA-CoA. O₂ was kept in the crystallographically observed position. All simulations were performed in an octahedral box of TIP3P water using periodic boundary conditions, with a size consistent with an O₂ concentration in the order of an O₂ saturated solution (5 mM; for comparison, an O₂ saturated buffer is ~1.2 mM). The net negative charge of the system was neutralized adding Na⁺ atoms. Details on how the system was minimized, heated and equilibrated are available in the ESI.† All production simulations were performed at 300 K, maintained using a Langevin thermostat with a 2 ps⁻¹ constant for temperature coupling. The SHAKE algorithm was used to keep bonds involving hydrogen atoms at their equilibrium length. Newton's equations were integrated using a 2 fs time step. Frames were collected every 5000 steps. 5 independent production simulations were performed, each approximately 100 ns long.

The free energy profile for O₂ migration in DpgC was computed using implicit ligand sampling (ILS) post-processing.³⁵ This method has previously been used to study migration of CO in heme proteins,^{46–48} and of O₂ in lipoyxygenase.⁴ Post-processing was performed on 3000 frames of one MD simulation of DpgC, and each subunit was processed individually in



parallel. ILS calculations were performed on a rectangular grid (0.5 Å resolution) using an O₂ molecule as probe.

Kinetic measurements

The kinetic parameters for DPA-CoA were determined using the DTNB [5,5'-dithiobis(2-nitrobenzoic acid)] reporter assay as described elsewhere^{24,33} with one modification: the concentration of enzyme used was 0.8 μM. Briefly, a freshly prepared solution of DTNB (1 mM), Tris-HCl (250 mM, pH = 7.5), and increasing concentrations of DPA-CoA were mixed with the enzyme (0.8 μM) at 24 °C (1 mL reaction volume). The reactions were monitored in a UV-vis spectrophotometer (Agilent) at 412 nm for 5 min. The observed initial rate of absorbance increase was converted to initial reaction velocity (V_0) using $\epsilon_{412\text{ nm}} = 13\,600\text{ M}^{-1}\text{ cm}^{-1}$. K_M^S and k_{cat}^S were determined by performing a non-linear regression to a plot of V_0 vs. [DPA-CoA].

The kinetic parameters for O₂ were determined as described elsewhere⁴⁹ using a fluorescence quenching based probe (Ocean Optics) in an airtight cuvette. All experiments were performed at 25 °C with continuous 1000 rpm stirring, using 83 μM DPA-CoA and varying concentrations of O₂. The reaction buffers were prepared by injecting different volumes of O₂-saturated buffer into the cuvette containing varying volumes of air-equilibrated buffer and DPA-CoA. The assay was initiated by injection of DpgC (1.67 μM) into the reaction vessel. Data was collected every 0.1 s. Initial velocities were determined from the initial change in oxygen concentration observed after accounting for non-enzymatic decrease in oxygen levels. Since the $K_M^{O_2}$ for DpgC is larger than the concentration of O₂ in a saturated solution,³³ it is not possible to obtain data above $K_M^{O_2}$ and it is more convenient to work under the assumption that $K_M^{O_2} \gg [O_2]$ and fit the data to a linear equation with slope $V_{\text{max}}^{O_2}/K_M^{O_2}$.

Xe-pressurized crystallography

DpgC-inhibitor co-complex crystals, obtained as described elsewhere,³³ were placed in a xenon pressure chamber (Hampden research) for ~15 minutes with pressures of 300 to 500 psi. The Xenon equilibrated crystals were flash frozen and diffraction data collected on the X12B beamline of NSLS at Brookhaven National Laboratory at a wavelength of 1.500 Å. The data was processed using the XDS program⁵⁰ in the space group $P2_1$ ($a = 139.1\text{ Å}$, $b = 171.0\text{ Å}$, $c = 156.0\text{ Å}$, $\beta = 90.02^\circ$). The structures were solved using the trimer from PDB 2NP9 as a starting model to find twelve monomers in the asymmetric unit (Matthews coefficient of $3.23\text{ Å}^3\text{ Da}^{-1}$ and 61.94% solvent). Xe sites were added manually based on anomalous difference electron density maps⁵¹ and the total structure was refined until $R_{\text{work}}/R_{\text{free}}$ converged to 0.17/0.20% using remlac⁵² (Table S9†). Structural figures were prepared using PyMOL⁵³ or VMD.⁵⁴

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