Multiple implications of an active site phenylalanine in the catalysis of aryl-alcohol oxidase

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Aryl-alcohol oxidase (AAO) has demonstrated to be an enzyme with a bright future ahead due to its biotechnological potential in deracemisation of chiral compounds, production of bioplastic precursors and other reactions of interest. Expanding our understanding on the AAO reaction mechanisms, through the investigation of its structure-function relationships, is crucial for its exploitation as an industrial biocatalyst. In this regard, previous computational studies suggested an active role for AAO Phe397 at the active-site entrance. This residue is located in a loop that partially covers the access to the cofactor forming a bottleneck together with two other aromatic residues. Kinetic and affinity spectroscopic studies, complemented with computational simulations using the recently developed adaptive-PELE technology, reveal that the Phe397 residue is important for product release and to help the substrates attain a catalytically relevant position within the active-site cavity. Moreover, removal of aromaticity at the 397 position impairs the oxygen-reduction activity of the enzyme. Experimental and computational findings agree very well in the timing of product release from AAO, and the simulations help to understand the experimental results. This highlights the potential of adaptive-PELE to provide answers to the questions raised by the empirical results in the study of enzyme mechanisms.

Elucidation of structure-function relationships is of key importance in the study of enzyme catalysis. Unveiling the mechanisms that lie behind the properties of a biocatalyst paves the way to widen its biotechnological applicability by means of enzyme engineering1. In this regard, aryl-alcohol oxidase (AAO) is an enzyme of biotechnological interest from the glucose-methanol-choline oxidase/dehydrogenase (GMC) superfamily. AAO catalyzes the oxidation of primary benzyl alcohols into their aldehyde counterparts using atmospheric O2 as co-substrate and yielding H2O2 as co-product2–4. In this way, it has shown its potential for the production of flavours and aromas, deracemisation of alcohol mixtures and the synthesis of precursors for the manufacturing of renewable polyesters5.

The ecophysiological role of AAO would be the supply of H2O2 to either ligninolytic peroxidases—which it acts synergistically with—or to trigger Fenton reactions during natural decay of lignocellulosic materials5–6. H2O2 production by AAO involves the redox-cycling of p-anisaldehyde7, a metabolite of AAO-producing species8–9. Its natural role as H2O2 producer may be exploited for biotechnological purposes by the development of enzyme cascades in which AAO and H2O2-consuming enzymes (peroxidases and peroxygenases) act concertedly. In recent years, the repertoire of AAO substrates has been enlarged by the discovery of new molecules the enzyme can oxidise, such as 5-hydroxymethylfurfural, 5-methoxymethylfurfural and their partially oxidised derivatives10,11, and secondary benzylic alcohols. Regarding the latter substrates, AAO stereoselectivity and activity was improved by means of computer-guided rational design that facilitated a more appropriate positioning of benzylic alcohols in the active site thanks to the removal of the Phe501 side chain12. AAO presents other potential applications in lignocellulose transformation and production of flavours and aromas13. Altogether, this variety of bioconversions renders AAO an enzyme with a bright future in biocatalysis.

The structure-function relationships of the model AAO from Pleurotus eryngii have been extensively investigated, and the roles of several residues important for catalysis have been elucidated14–17. The catalytic pocket of P. eryngii AAO, located near the flavin moiety of its FAD cofactor, is shielded from the outer environment by a triad

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of aromatic residues —Tyr92, Phe501 and Phe397— that form a hydrophobic bottleneck affecting alcohol and oxygen migrations into AAO active-site\textsuperscript{18,19}. The roles of Tyr92 and Phe501 have been unveiled and are involved in: (i) establishing aromatic stacking interactions that guide the alcohol substrate to a catalytically competent configuration\textsuperscript{13}; and (ii) compressing the active site to promote the reactivity with O\textsubscript{2}, respectively. This residue is located in a loop characteristic of the AAO family (residues 395–406 in \textit{P. eryngii} AAO), which partially covers the access to the active site\textsuperscript{20} (Fig. 1). Phe397 has been proposed to act as a barrier that prevents the free diffusion of molecules —substrates and/or products— in and out of the catalytic pocket protecting the flavin environment of AAO. Previous computational studies of substrate migrations into the active site —using the PELE algorithm\textsuperscript{21}— suggested that Phe397 swung along with the alcohol substrate, helping it to reach the catalytic pocket \textsuperscript{19} (www.youtube.com/watch?v=CqSDn5OmagI). These results have encouraged the investigation of the role of Phe397 in AAO catalysis.

In this work, kinetic, ligand binding and diffusion studies on AAO and several mutated variants have been performed to enlighten our understanding on the role of Phe397 in catalysis. Moreover, molecular dynamics and new ligand diffusion computational studies using adaptive-PELE —a new version of the PELE algorithm that avoids the metastability of the ligands, thus saving number of processors and computation time\textsuperscript{22}— have been employed to shed light on the catalytic implications of substituting Phe397 in AAO. Experimental findings and computational results agree very nicely and prove the applicability of the new adaptive-PELE to modelling the dynamic nature of protein-ligand interactions and its ability to unveil the mechanisms underlying binding in complex systems.

Results
Spectral properties and steady-state kinetics of AAO and its Phe397 variants. Wild-type recombinant AAO (hereinafter native AAO) and its F397A, F397L, F397Y and F397W variants were purified as holoproteins, after \textit{Escherichia coli} expression and \textit{in vitro} activation. The UV-visible spectra of all variants showed the typical flavin bands around 460 and 385 nm and shoulder around 500 nm (data not shown), indicating proper folding around the cofactor. 280 nm/460 nm absorbance ratios around 10–11 showed that the FAD cofactor was in the oxidized state and correctly incorporated into the protein for all variants.

The redox state of the cofactor during turnover was investigated by following the spectra of the AAO variants during oxidation of \textit{p}-methoxybenzyl alcohol (initially saturated with air) until the enzyme was completely reduced. In the cases of native AAO and the majority of the variants —F397W, F397A and F397L— the most abundant species during turnover (initial reaction phase) is the oxidized enzyme, whereas the reduced species is the predominant for F397Y (Fig. 2).

Bi-substrate steady-state kinetics —measured as \textit{p}-anisaldehyde production from \textit{p}-methoxybenzyl alcohol at different O\textsubscript{2} concentrations— revealed remarkable differences among native AAO and its F397 variants (Table 1). First, in all four variants, contrary to the native AAO\textsuperscript{26}, the kinetics best fitted equation (1) describing a ping-pong mechanism, as revealed by the Hanes-Woolf plots of its bi-substrate kinetics (Supplementary Fig. S1). Then, all variants, with the exception of F397L, showed 2–3 fold lower turnover rates (\(k_{\text{cat}}\) values) than the native enzyme. Regarding affinity for the alcohol substrate, the F397Y variant showed the same Michaelis constant (\(K_{m(\text{Al})}\) as

![Figure 1. Surface loop and aromatic residues limiting access to the AAO FAD. (a) Area of channel opening on the AAO surface, with the 395–406 loop in transparent light blue. (b) Semitransparent surface showing the aromatic gate-keeping Tyr92, Phe397 and Phe501 aromatic residues (sticks in olive green), loop (green) and its contiguous \(\beta\)-strand residues (pale yellow cartoon), the catalytic His502 (grey) and FAD (grey), with one \textit{p}-anisic acid molecule at the active site. From AAO: \textit{p}-anisic acid structure (PDB 5OC1)\textsuperscript{17}.](image-url)
the native protein, while the F397A, F397L and F397W substitutions increased $K_{m(Al)}$ by ~2-, 9- and 11-fold, respectively. Consequently, the F397Y, F397A, F397L and F397W variants were ~3-, 4-, 10- and 21-fold less efficient ($k_{cat}/K_{m}$) oxidizing the alcohol substrate compared to the native AAO. Regarding the affinity for O$_2$, all the variants, with the exception of F397A, showed increased affinity (lower $K_{m(ox)}$) with regard to the native protein. Thus, F397Y, F397L and F397W were 1.4-, 1.6-, and 2-fold more efficient using O$_2$ as an electron acceptor than the native protein. Finally, steady-state constants calculated from anisaldehyde release (as in previous experiments) were compared with those obtained from H$_2$O$_2$ release (in additional kinetic analyses under atmospheric O$_2$ saturation). In all variants, $k_{cat}$ and $K_{m(Al)}$ tended to be almost identical using both approaches (Supplementary Table S1), as it had been previously reported for native AAO$^{23}$.

Rapid kinetics of the two half-reactions for the Phe397 variants. In the light of the above results, the reductive and oxidative half-reactions of the F397 variants were analyzed to unveil the rate-limiting step during catalysis. The spectra collected during the reductive half-reactions of the flavin, in agreement with the previously reported hydride transfer reaction for the native AAO$^{16}$, global analyses of the spectral evolution were fitted to a one-step model (A $\rightarrow$ B) in all cases (Fig. 3). The values of the observed rate constants ($k_{red}$) at different substrate concentrations exhibited a hyperbolic dependence on the alcohol concentration (Supplementary Fig. S2) that allowed the determination of the reduction rate constant ($k_{red}$) and the dissociation constant ($K_{d(Al)}$) upon fitting to equation (3). For native AAO (spectral changes not shown in Fig. 3) and the F397A and F397L variants, the $k_{red}$ values (Table 2) were of the same range of the previously determined $k_{cat}$ values (Table 1) indicating that the reductive half-reaction is the rate-limiting step in catalysis. Nevertheless, variants F397Y and F397W showed $k_{red}$ values 3- and 2-fold higher than the respective turnover rates, suggesting that reductive half-reaction is not the limiting step in these variants. For all the F397 variants, $K_{d(Al)}$ values were similar to the $K_{m(Al)}$ estimated under steady-state conditions.

The oxidative half-reactions of the four Phe397 variants, and native AAO (not shown), fitted two-step model equations (A $\rightarrow$ B $\rightarrow$ C) describing a biphasic pattern (Fig. 4) where the first phase accounts for more than...
70–80% of the total amplitude. For the F397Y, F397L and F397A variants, this \( k_\text{obsA} \) was linearly dependent on \( O_2 \) concentration, allowing the determination of a second-order rate constant \( (k_\text{ox}/K_{d(\text{ox})}) \) that was similar, 2– or 10-fold slower than that of native AAO, respectively (Table 2). For the F397L and F397A variants, a reverse rate constant was observed, \( k_\text{rev} \sim 15 \text{s}^{-1} \), which corresponds to the intercept of the y-axis (Supplementary Fig. S3A). On the contrary, the \( k_\text{obsA} \) for F397W showed a hyperbolic dependence on \( O_2 \) concentration that fitted equation (5) and describes an encounter complex of a reduced flavin enzyme with \( O_2 \) followed by flavin reoxidation (Supplementary Fig. S3B). The estimated transient-state second order rate constant, \( k_\text{ox}/K_{d(\text{ox})} \sim 689 \text{s}^{-1}\text{mM}^{-1} \) (Table 2 footnote) agreed with the catalytic efficiency determined under steady-state conditions (Table 1).
The second phase for all variants was too slow to be relevant for catalysis ($k_{\text{obs}} < 7 \text{ s}^{-1}$), although it was independent of O$_2$ concentration for the F397Y and F397W variants and dependent on it in the case of the F397L and F397A variants (Supplementary Fig. S3A, inset). The O$_2$-independent slow phase in native AAO reoxidation had been previously attributed to the presence of damaged protein ensuing the stopped-flow experiments. The O$_2$-independent slow phase in native AAO reoxidation had been previously attributed to the presence of damaged protein ensuing the stopped-flow experiments.

**Studies on AAO: $p$-anisic acid complex formation and dissociation.** The differences between the $k_{\text{cat}}$ and $k_{\text{in}}$ values for both F397Y and F397W encouraged us to investigate whether product release has an effect on turnover. The study could not be carried out with $p$-anisaldehyde because the formation of the enzyme-aldehyde complex is too rapid, which prevents its detection by the stopped flow equipment. Therefore, this study (Fig. 5) was performed with the final product, $p$-anisic acid, whose crystallographic complex with AAO was recently solved (PDB 5OC1). The rate constants for complex formation ($k_{\text{cat}}$) and dissociation ($k_{\text{in}}$) of native AAO and the F397Y and F397W variants are shown in Table 3. The $k_{\text{cat}}$ values for F397Y and F397W were 2-fold and 2 orders of magnitude lower than that of the native enzyme, respectively. Differences in $k_{\text{in}}$, were also observed, since the $k_{\text{in}}$, values for these variants were at least 10-fold slower than that of the native AAO. The fast diffusion of the ligand in and out the active site of the F397L and F397A variants most likely impaired estimation of these parameters. The $K_{\text{d(Ac)}}$ values calculated from the corresponding $k_{\text{in}}$/$k_{\text{cat}}$ ratios for native AAO and the F397Y and F397W variants (150, 31 and 271 µM respectively) agree well with the values obtained by differential spectrophotometry (Supplementary Fig. S4 and Table 3). F397Y showed the smallest $K_{\text{d(Ac)}}$, which means that it is the variant that more tightly binds $p$-anisic acid. On the contrary, the $K_{\text{d(Ac)}}$ for the F397L and F397A variants could not be estimated by differential spectrophotometry since saturation could not be attained in agreement with the above results.

**AAO molecular dynamics and ligand diffusion simulations.** MD simulations were carried out to study the dynamical behaviour of the five protein systems (native enzyme and its Phe397 variants). Loops are the secondary structures that most likely adopt different conformational structures. Hence, the motions of the loops Gln395-Thr406 and Ser89-Met95, where two of the main gate residues—Phe397 and Tyr92—are located, were investigated. However, the root-mean-square-deviation values obtained, as a function of time, were low (data not shown) indicating that there are only moderate displacements of these secondary structures along the dynamics of the protein.
Despite the small loop movements, a closer look at the distances between the gate residues indicates that the F397Y and F397W substitutions induce significantly shorter values (Supplementary Fig. S5A). Moreover, inspection of the protein structures shows that the side chains of the F397Y and F397W variants establish hydrogen bonds with Tyr92 with a frequency of 20% and 5%, respectively (Supplementary Fig. S5B), while the aliphatic variants and the native enzyme cannot develop such hydrogen bonds resulting in larger distances between gate residues. Altogether, the MD data suggests local constraints induced by the tyrosine and tryptophan substitutions in the gate region.

The diffusion of products — $p$-anisaldehyde and $p$-anisic acid — and reactants — $p$-methoxybenzyl alcohol and $O_2$— was also investigated with adaptive-PELE to explain the changes in kinetics observed for the mutants.

The trajectory analysis of the acidic product in the aromatic variants clearly indicates a stabilization of the local

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**Table 3.** Dissociation constants, and formation/dissociation rates of $p$-anisic acid complexes. The dissociation constants ($K_{d(Ac)}$) of $p$-anisic acid complexes with AAO and its Phe397 variants were measured by differential spectroscopy, while the complex formation ($k_{for}$) and dissociation ($k_{dis}$) rates were measured by transient-state kinetics spectroscopy. All experiments were performed in 50 mM sodium phosphate (pH 6.0) at 12 °C. nd, not determined (too fast complex formation and dissociation). Means and standard deviations from the fit to equations (7) or (6). All data were measured by triplicates.

|          | $K_{d(Ac)}$ (µM) | $k_{for}$ (s$^{-1}$·mM$^{-1}$) | $k_{dis}$ (s$^{-1}$) |
|----------|-----------------|-----------------------------|-----------------|
| AAO      | 170 ± 5         | 18.0 ± 0.3                  | 2.7 ± 0.17      |
| F397Y    | 25 ± 2          | 7.7 ± 0.1                   | 0.24 ± 0.04     |
| F397W    | 200 ± 12        | 0.70 ± 0.01                 | 0.19 ± 0.01     |
| F397A    | nd              | nd                          | nd              |
| F397L    | nd              | nd                          | nd              |

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**Figure 5.** Spectral changes during formation of AAO-$p$-anisic acid complex. (a) Native AAO, mixed with 0.5 mM of the ligand at 0.003, 0.01, 0.02, 0.03, 0.04, 0.05, 0.065, 0.085, 0.1, 0.13, 0.15, 0.2 and 0.4 s. (b) F397Y, mixed with 0.65 mM of $p$-anisic acid measured after 0.005, 0.05, 1.0, 2.0, 3.0, 4.0, 5.0, 6.0, 7.0, 8.0, 9.0, 1.2, 3 and 4 s, and (c) F397W mixed with $p$-anisic acid 1 mM recorded at 0.005, 0.02, 0.04, 0.05, 0.07, 0.1, 0.2, 0.4, 0.5, 0.7, 1.2 and 5 s. Insets show disappearance of initial species A and formation of final species B. The rate constant obtained from the global fitting of the spectral changes is shown. (d) Fits of the obtained $k_{obs}$ for each variant and ligand concentration to equation (6). Native AAO (filled circles), and F397Y (open circles) and F397W (inverted triangles) variants.
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Figure 6. Boxplots of PELE steps required for ligand release from AAO and its Phe397 variants. (a) p-Anisic acid. (b) p-Anisaldehyde. Number of PELE steps is directly related to time. Grey, native AAO; blue, F397W; red, F397Y; yellow, F397A; and light green, F397L variants. Boxes contain 50% of the results (two quartiles), horizontal lines inside them indicate the mean value; while the upper and lower whiskers (vertical lines) contain the remaining quartiles (50%). The overall height of boxes (whiskers included) is indicative of the spread of the results. Isolated points represent results significantly different from the rest of data values.

Discussion

Phe397 is situated at a loop characteristic of the AAO family, in a region prone to harbouring insertions and deletions amongst the members of the GMC superfamily. Representatives of this superfamily are monomeric or multimeric proteins, with the entrance to their active site partially covered by the adjacent monomer in the second case. Given their monomeric nature, AAOs and a few other GMCs have developed loop structures to control diffusion of molecules into their active sites. Among them, choline oxidase25, cholesterol oxidase 26 and cellobiose dehydrogenase27 are notable examples, whose insertions are even longer than the AAO insertion. The latter insertion partially forms the 395–406 loop in P. eryngii AAO and encloses the active-site cavity from the outer environment (Fig. 1)18. Previous computational studies hinted that Phe397 oscillates with the substrate as a mechanism of gating28, similarly to what has been reported for phenylalanine residues in P450 enzymes 28. A homologous phenylalanine residue is conserved in 50% of all the putative AAO sequences from basidiomycete genomes available at JGI (https://jgi.doe.gov). The multiple roles of this phenylalanine residue in catalysis, as revealed by studies on the P. eryngii model AAO, are discussed below.

Phe397 favours the correct positioning of the alcohol and its oxidation at the active site, as evidenced by the steep decrease of catalytic efficiency in the mutated variants. Such effect is most noticeable in F397W, which shows the lowest $k_{cat}/K_m (Al)$ value due to: i) the constrained distance between gate residues, and ii) the formation of a H bond with its reactive hydroxyl group that disrupts the substrate's catalytic poses. The introduction of a bulkier residue at the Phe501 and Tyr92 positions of AAO also produced similar steric hindrances for catalysis13,14,17. Remarkably, the drop of the catalytic efficiency in F397Y is due to its small turn over rates, as affinity for alcohol ($K_{d(Al)}$) and $K_{d(Al)}$ remains invariable in this case. Furthermore, the reduction rates being higher than turnover in F397Y and F397W, indicates that, unlike native enzyme, the reductive half-reaction is not the rate-determining step in these two variants, as discussed below.

Phe397 also plays a role in flavin reoxidation by either compressing the active site to reduce the free diffusion of O$_2$, or altering its redox environment. Evidence comes from the low oxidation rates estimated for the aliphatic mutants, in contrast to the ones obtained for the aromatic variants and the native AAO, which possess bulkier residues mentioned above.

Furthermore, the distances among any of the atoms of the O$_2$ substrate and both the C4a locus of reduced FAD and the N$_{e2}$ of His502 (acting as a catalytic base in the reductive half reaction) were calculated for all the variants. Results suggest that the catalytic distances are similar in the native enzyme and variants, except for F397W that shows higher population at favourable —that is, shorter— distances (Fig. 7).

Finally, the energetic profiles of the $p$-methoxybenzyl alcohol catalytic distances (hydroxyl-to-His502 and proR-H-to-flavin) in all protein systems were studied, as a measure of the binding effectiveness of the alcohol in the active centre. The energy landscape shows an additional energetic favourable region only for the F397W. In this new minimum, the side chain of the tryptophan interacts via hydrogen bond to the hydroxyl group of the alcohol substrate, resulting in an inefficient catalytic pose (Supplementary Fig. S6). This is in agreement with the experimental values, where the $K_{d(Al)}$ and $K_{d(Al)}$ are almost 11-fold higher for the F397W variant compared to the native protein.
residues that constrain the catalytic pocket. However, the oxidative half-reaction does not limit the catalytic cycle in the aliphatic variants, as $k_{\text{cat}}$ and $k_{\text{red}}$ values are similar and the constants measured as $\text{H}_2\text{O}_2$ release are identical as those measured by aldehyde formation. Alteration of the FAD environment and electrochemistry might be involved in the decreased efficiency for oxidation of the F397A cofactor. Regarding the F397Y and F397W variants, they show high $k_{\text{ox}}$ similar to the native protein. In the case of F397W variant, the obtained $k_{\text{ox}}$ and $k_{\text{red}}$ are similar indicating that the reductive and oxidative half-reactions are almost balanced, which agrees with its redox state during turnover. Computational simulations of F397W reoxidation revealed that withdrawal of His502 due to the presence of the tryptophan increases the ability of this variant for properly positioning the oxygen molecule during flavin reoxidation, bringing $O_2$ and C4a closer than in the native enzyme (Fig. 7). However, the introduction of this bulkier residue could be hindering the oxygen diffusion into the active-site 19 and thus causing the unusual saturation behaviour observed for this variant with increasing oxygen concentration. This saturation profile has been observed in cholesterol oxidase from *Brevibacterium sterolicum* related to the interconversion between open and closed channel enzyme conformations. This event regulated the oxygen diffusion and constituted the rate-limiting step preceding flavin reoxidation39,30. Regarding F397A and F397L, they display slower reoxidation rates probably due to the absence of an aromatic residue in that position. Moreover, both variants display slow phases that are dependent on $O_2$ concentration, that could reveal two different kinetic processes, the H and H$^+$ transfers recently reported to take place in separate kinetic steps in native AAO 24.

Finally, Phe397 also plays a role in controlling the product release from the active site, with this catalytic step limiting the turnover of the F397Y and F397W variants, while the F397A and F397L variants (and native AAO) let the substrate and product diffuse easily in and out of the active site. Furthermore, ligand migration studies with both $p$-anisic acid and $p$-anisaldehyde indicate that product release is delayed due to the reduced gated space—which forces the aromatic side chains to be displaced—and the hydrogen bond between the tyrosine or tryptophan and Tyr92 in the aromatic variants. The native enzyme also requires longer times for product release because its side chain must move to let it out, although there is no hydrogen bonding between the Phe397 and Tyr92. This is in agreement with the 10-fold lower $k_{\text{cat}}$ for the aromatic variants than for native AAO. Flavoenzymes with catalytic cycles limited or partially limited by the product release have been reported, such as D-amino acid oxidase31, cyclohexanone monooxygenase32 or nitroalkane oxidase33. The case of amadoriase I is also relevant, since

**Figure 7.** $O_2$ diffusion in native AAO and its F397W variant. (a) Native and F397W final catalytic poses (CPK-colored molecules with grey and cyan carbon atoms, respectively) showing catalytic distances (in Å) to FAD C4a and His502. (b and c) Frequency of $O_2$ poses as a function of the distances from $O_2$ to His502 and C4a of native and the F397W variant, respectively.
its catalysis is partially limited by the product release, although the oxidation of the enzyme takes place through an ordered ternary complex of enzyme, product and O₂. 

In conclusion, Phe397 plays a central role in the catalysis of P. eryngii AAO, which is reinforced by its high conservation among other sequences of characterised and putative AAOs from fungal genomes. As drawn from the experimental and computational results, this residue supports both half-reactions of the enzyme. During the reductive reaction, it helps the substrate attain a catalytically relevant position and facilitates product release from the active site. Analysis of the oxidative reaction suggests that the presence of an aromatic residue at this position is important to modulate the flavin environment and to reduce the space in the active site enabling the enzyme to reduce O₂ to H₂O₂. Experimental and computational results complement each other well and, thus, highlight the suitability of the recently developed adaptive-PELE for the study of protein binding to different ligand types.

**Methods**

**Reagents.** Glucose oxidase type VII from Aspergillus niger, glucose, p-methoxybenzyl alcohol and p-anisic acid were purchased from Sigma-Aldrich.

**Enzyme expression and purification.** Native AAO from P. eryngii (GenBank accession number AF064069) and its mutated variants F397A, F397Y, F397W and F397L were heterologously expressed in E. coli W3110 as recombiant proteins using the pFLAG1 vector. The above-mentioned AAO variants were prepared using the QuickChange® site-directed mutagenesis kit.

PCR reactions were run with the following oligonucleotides harbouring the desired mutations (underlined nucleotide/s in bold triplets indicated below) and the AAO gene inserted in the vector as a template: (i) F397Y: 5'-CTTTCACCAATGTTACCCACCGTATCTCCTG-3', (ii) F397A: 5'-CTTTTCACCAATGTTACCCACCGTATCTCCTG-3', (iii) F397W: 5'-CTTTTCACCAATGTTACCCACCGTATCTCCTG-3', and (iv) F397L: 5'-CTTTTCACCAATGTTACCCACCGTATCTCCTG3', along with their reverse complementary counterparts. The AAO variants were expressed as inclusion bodies, subsequently purified by anion-exchange chromatography as previously described.

Molar absorbance coefficients of the variants were calculated by heat denaturation of the proteins and determination of the FAD released. The coefficients for native AAO and the F397Y, F397A, F397W and F397L variants are ε₄₆₂ nm = 11050 M⁻¹ cm⁻¹, ε₄₆₂ nm = 10415 M⁻¹ cm⁻¹, ε₄₃₉ nm = 9432 M⁻¹ cm⁻¹, ε₄₆₂ nm = 9952 M⁻¹ cm⁻¹ (at 462 nm), and ε₄₆₂ nm = 12492 M⁻¹ cm⁻¹, respectively.

**Steady-state kinetics.** Bi-substrate kinetics of the four AAO variants were measured recording p-anisaldehyde production (ΔΔε₅₆₃ = 16950 M⁻¹ cm⁻¹), varying both the concentrations of p-methoxybenzyl alcohol and O₂ in a spectrophotometer. Reactions were performed in screw-cap cuvettes, where different O₂ concentrations in 0.1 M phosphate at pH 6.0, were attained by bubbling several O₂/N₂ mixtures inside a thermostatic bath at 12°C. Reactions were triggered by the addition of the alcohol substrate (5–500 μM) and the enzyme (3–5 nM) using microsyringes. The spectrophotometer was maintained at the desired temperature owing to the use of a thermostatic bath.

Initial rates were obtained from the linear phase of the aldehyde production and were calculated as the change in absorbance over time. Kinetic constants were estimated by fitting the kₘ to equation (1) describing a ping-pong mechanism:

$$\frac{\nu}{e} = \frac{k_{cat}AB}{K_{m(Al)}B + K_{m(Ox)}A + AB}$$

(1)

where ν stands for the initial velocity, e is the enzyme concentration, kₖₐₜ is the catalytic constant, A stands for the alcohol concentration, B represents O₂ concentration and Kₘ(Al) and Kₘ(Ox) are the Michaelis constants for p-methoxybenzyl alcohol and O₂, respectively.

Kinetic constants estimated as p-anisaldehyde and H₂O₂ release were compared under atmospheric O₂ saturation conditions, at 25°C. p-Anisaldehyde was measured as detailed above, while H₂O₂ release was estimated by coupling the reaction of a horseradish peroxidase (6 U·mL⁻¹) and AmplexRed® (60 nM), which uses the H₂O₂ produced by AAO to give coloured resorufin (ΔΔε₅₆₃ = 52000 M⁻¹ cm⁻¹). Catalytic constants were estimated by fitting to a Michaelis-Menten equation for one substrate in both cases (equation (2)):

$$\frac{\nu}{e} = \frac{k_{cat}A}{K_{m(Al)} + A}$$

(2)

**Stopped-flow measurements.** Experiments were performed using a stopped-flow spectrophotometer from Applied Photophysics Ltd. model SX17.NV.

Enzyme-monitored turnover experiments were carried out by mixing AAO (~10 μM) with exceedingly saturating p-methoxybenzyl alcohol concentrations (at least 10-fold the Kₘ for each of the variants), under air-saturated conditions, at 25°C. Spectral evolution of the enzymes during redox turnover was recorded between 350 and 700 nm with a photodiode array (FDA) detector.

Studies on the reductive half-reaction were performed upon mixing AAO with increasing concentrations of p-methoxybenzyl alcohol (0.018–0.6 mM) under anaerobic conditions. The stopped-flow apparatus was made aerobic by flushing sodium dithionite through the system, which was then rinsed out with O₂-free buffer. All buffers, substrates and the enzyme were poured into glass tonometers that were subsequently subjected to 20–25 cycles.
of evacuation and argon (Ar) flushing in order to remove O$_2$. To ensure anaerobiosis, glucose (10 mM) and glucose oxidase (10 U·mL$^{-1}$) were added after some vacuum-Ar cycles. Measurements were recorded using the PDA detector, in 50 mM sodium phosphate (pH 6.0) at 12 °C. Observed rate constants ($k_{\text{obs}}$) were calculated by global fitting of the spectra with Pro-K software to a single exponential equation. Those averaged $k_{\text{obs}}$ at each substrate concentration were then non-linearly fitted to equation (3), describing hyperbolic substrate dependence of $k_{\text{obs}}$:

$$k_{\text{obs}} = \frac{k_{\text{red}}A}{K_{d(\text{Al})} + A} + k_{\text{rev}}$$  \hspace{1cm} (3)

where $k_{\text{red}}$ and $k_{\text{rev}}$ represent the reduction rate constant at infinite substrate concentration and its reverse reaction, respectively; $A$ stands for the alcohol concentration; and $K_{d(\text{Al})}$ is the dissociation constant.

The oxidative half-reaction was investigated by mixing reduced AAO with increasing O$_2$ concentrations. Procedures were as explained above for the reductive half-reaction, except that, in this case, AAO and glucose were put into a tonometer bearing a side-arm, where p-methoxybenzyl alcohol (1.3-fold the concentration of AAO) was poured along with glucose oxidase. After the required vacuum-Ar cycles, enzyme and substrate were mixed before being mounted onto the stopped-flow equipment. Reactions were measured with both the PDA and the monochromator detectors at 12 °C. $k_{\text{obs}}$ were obtained by either global fitting of the spectra or fitting the monochromator traces to exponential equations describing two-step and three-step processes. Fitting averaged $k_{\text{obs}}$ either to equation (4) that describes a linear dependence on O$_2$ concentration or equation (5) that describes hyperbolic saturation with increasing O$_2$ concentration allowed the estimation of the apparent second-order rate constant for reoxidation ($k_{\text{app}}$) and the first-order rate ($k_{\text{fot}}$) and second-order constants for reoxidation ($k_{\text{fot}}/K_{d(\text{ox})}$), respectively:

$$k_{\text{obs}} = k_{\text{fot}} + k_{\text{dis}}$$  \hspace{1cm} (4)

Estimation of the rates of the AAO:p-anisic acid complex formation and dissociation were performed by analyzing spectral changes upon mixing enzyme (~10 µM) with different concentrations of the ligand (0.04–2 mM) at 12 °C. Data were globally fitted to an equation describing a one-step process. The obtained $k_{\text{obs}}$ were linearly dependent on the ligand concentration and were, thus, fitted to equation (6):

$$k_{\text{obs}} = k_{\text{fot}}L + k_{\text{dis}}$$  \hspace{1cm} (6)

in which $k_{\text{dis}}$ stands for the second-order rate constant for the complex formation; $L$ represents the ligand concentration, and $k_{\text{dis}}$ is the rate constant for the complex dissociation.

**Spectral characterization of the AAO:p-anisic acid complex.** The affinity of the native AAO and its variants for p-anisic acid was assessed by titrating the enzyme with increasing concentrations of the ligand in 50 mM sodium phosphate (pH 6.0) at 12 °C. Spectral changes were recorded using a spectrophotometer and their magnitude upon complex formation was fitted to equation (7), which accounts for a 1:1 stoichiometry, as a function of p-anisic acid concentration:

$$\Delta A = \frac{\Delta \varepsilon(E + L + K_d) - \Delta \varepsilon(E + L + K_d)^2 - 4EL}{2}$$  \hspace{1cm} (7)

in which $\Delta A$ accounts for the observed change in absorbance; $\Delta \varepsilon$ represents the maximal absorption difference in each of the spectra; $K_d$ is the dissociation constant; and $E$ and $L$, the enzyme and p-anisic acid concentrations.

**Molecular dynamics (MD) studies.** The protonation states of the titratable amino acids present at the AAO crystal structure (PDB 3FIM) were set up using PROPKA$^{46}$ of Protein Preparation Wizard$^{47}$ and checked with the H++ server$^{48}$. The selected mutations (F397W, F397Y, F397A and F397L) were introduced and all 3D structures were minimized with Prime$^{39,40}$. The overall systems were solvated and neutralized using Desmond$^{41}$ with the SPC water model and 0.15 M of NaCl.

50 ns of MD simulation were performed with Desmond$^{41}$ for each protein structure using the default Desmond’s protocol. The temperature was regulated with the Nosé-Hoover chain thermostat$^{42–44}$ with a relaxation time of 1 ps. The pressure was controlled with the Martyna-Tobias-Klein barostat$^{45}$ with isotropic coupling and a relaxation time of 2 ps. RESP integrator$^{46,47}$ was used with bonded, near and far time steps of 2.0, 2.0 and 6.0 ps, respectively. Furthermore, a 9 Å cut-off was employed for nonbonded interactions with the smooth particle mesh Ewald method$^{47}$.

**Ligand diffusion.** The minimized models described above were also used to study the ligand diffusion. The acid product was manually docked inside the cavity, next to Phe397 and Tyr92, while reactants were docked in their catalytic position. The FAD cofactor was protonated to its semiquinone state for the O$_2$ simulations, while it remained in its quinone state for the acid, aldehyde and alcohol diffusion studies.

The FAD cofactor was optimized with QM/MM at the M06/6–31G* level of theory using Qsite$^{48}$ and the atomic charges were obtained from the electrostatic potential (ESP). Ligand geometries were optimized at the
same level of theory in gas phase and re-optimized using the PBF implicit solvent with the Jaguar software. The ligands were parameterized according to the OPLS 2005 force field, maintaining the ESP charges, and a rotamer library was build using Macromodel.

Ligand diffusion was modelled with the recent adaptive-PELE technique22, using an anisotropic network model (ANM) backbone perturbation30 applied to all Cα atoms, side chains re-sampling within 6 Å of the ligands, and a full energy minimization in each PELE step. The products were randomly translated and rotated within a spherical box of 22 Å of its initial position. The O₂ and the alcohol reactants were randomly roto-translated within a spherical box of 3 Å and 7 Å of its centre of mass initial position, respectively, to inquire about their poses in the active centre. Adaptive-PELE simulations involved ~50 epochs of 4 PELE steps. All simulations were performed using 32 processors. The O₂ simulations were executed during 72 h, while the other simulations were performed during 23 hours.

Data availability statement. All data generated or analysed during this study are included in this published article (and its Supplementary Information files).

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Author Contributions
J.C. performed the experiments and wrote the experimental sections of the paper. P.A.-R. and F.S. carried out the computational simulations and wrote the computational sections of the manuscript. A.T.M., P.F. and M.M. designed and supervised the experimental studies. V.G. designed and supervised the computational studies. A.T.M., P.F., V.G. and M.M. supervised the discussion of results and main conclusions. All authors revised and approved the submitted manuscript.

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