Evolutionary History of a Specialized P450 Propane Monooxygenase

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The evolutionary pressures that shaped the specificity and catalytic efficiency of enzymes can only be speculated. While directed evolution experiments show that new functions can be acquired under positive selection with few mutations, the role of negative selection in eliminating undesired activities and achieving high specificity remains unclear. Here we examine intermediates along the ‘lineage’ from a naturally occurring C₁₂–C₂₀ fatty acid hydroxylase (P₄₅₀BM₃) to a laboratory-evolved P₄₅₀ propane monooxygenase (P₄₅₀PMO) having 20 heme domain substitutions compared to P₄₅₀BM₃. Biochemical, crystallographic, and computational analyses show that a minimal perturbation of the P₄₅₀BM₃ fold and substrate-binding pocket accompanies a significant broadening of enzyme substrate range and the emergence of propane activity. In contrast, refinement of the enzyme catalytic efficiency for propane oxidation (∼9000-fold increase in $k_{cat}/K_m$) involves profound reshaping and partitioning of the substrate access pathway. Remodeling of the substrate-recognition mechanisms ultimately results in remarkable narrowing of the substrate profile around propane and enables the acquisition of a basal iodomethane dehalogenase activity as yet unknown in natural alkane monooxygenases. A highly destabilizing L₁₈₈P substitution in a region of the enzyme that undergoes a large conformational change during catalysis plays an important role in adaptation to the gaseous alkane. This work demonstrates that positive selection alone is sufficient to completely respecialize the cytochrome P₄₅₀ for function on a nonnative substrate.

Keywords: alkane oxidation; protein evolution; directed evolution; propane monooxygenase; cytochrome P₄₅₀ BM3

Introduction

Organisms can adapt to changing environments by acquiring alternative biosynthetic pathways or detoxification mechanisms upon discovery and subsequent optimization of new enzyme catalysts. Despite fundamental contributions from genetic and structural analyses,¹,² our understanding of the forces driving the diversification and specialization of enzymes to occupy new functional niches remains limited.

Directed evolution experiments have demonstrated that a small number of mutations can alter enzyme specificity and enhance activity toward poor substrates.³⁻⁸ Catalytic promiscuity and the ease with which these side activities can be improved provide a mechanism for enzyme diversification in nature.⁹,¹⁰ Although there are cases in which having a broad substrate range is linked to the physiological role of the enzyme (e.g., the ability of certain P₄₅₀s to metabolize a wide array of xenobiotics),¹¹⁻¹⁴ many enzymes show exceptional power to discriminate among similar substrates. How this specialization occurs is largely unknown. In the laboratory, newly specialized enzyme variants have been obtained by applying positive and
negative selection pressures that reinforce the desired phenotype while removing unfavorable side activities. Negative selection may guide natural enzyme evolution in those cases where a promiscuous activity has a negative impact on viability, leading, for example, to accumulation of toxic intermediates or the waste of cellular resources. However, negative selection may not be required to achieve high specificity. It has been postulated that ‘generalist’ enzymes may have to trade off high activity on any one substrate for the ability to recognize and react with a wide range of substrates. ‘Specialist’ enzymes could thus be the natural result of maximizing activity on one particular substrate and not require negative selection to remove other activities. Specialization under positive selection alone, however, has not yet been demonstrated experimentally.

A large family of heme-containing monoxygenases comprising more than 7000 known members, the cytochrome P450s provide an extraordinary example of how a conserved structural fold diversified over millions of years to catalyze oxygenation reactions on widely different substrates. In addition to their key roles in the breakdown of xenobiotics, cytochrome P450s catalyze important steps in the biosynthesis of flavonoids, alkaloids, terpenes, sterols, and other natural products. Unlike the xenobiotic-oxidizing P450s, these latter, anabolic enzymes tend to display narrow substrate preference. We have recently reported the engineering of a proficient P450 propane monoxygenase (P450PMO)Applying directed mutagenesis. P450 PMO is highly active, supporting 33,400 turnovers on propane (versus 45,800 for P450PMOR2) with a coupling efficiency of 98% for P450PMOR2). The series begins with the octane-hydroxylating variant 139-3 and includes subsequent variants that show at least a twofold increase in propane total turnover number (TTN) and/or >10% improvement in coupling efficiency for propane oxidation compared to their predecessors. P450PMO,R2 carries a reductase domain mutation (D698G) beneficial to propane oxidation. To facilitate comparison with wild-type P450BM3 and other variants, we generated P450PMO,R2 with Gly698 reverted to Asp by site-directed mutagenesis. P450PMO,R2 is highly active, supporting 33,400 turnovers on propane (versus 45,800 for P450PMO,R2) with a coupling efficiency of 93% (versus 98% for P450PMO,R2). The mutagenesis/screening methods used for the isolation of the selected variants together with their amino acid substitutions are summarized in Table 1.

### Results

The ‘evolutionary lineage’ connecting P450BM3 to P450PMO was reconstructed through eight variants. The series begins with the octane-hydroxylating variant 139-3 and includes subsequent variants that show at least a twofold increase in propane total turnover number (TTN) and/or >10% improvement in coupling efficiency for propane oxidation compared to their predecessors.

#### Biochemical analysis of P450PMO and its evolutionary precursors

Changes in the catalytic parameters and stabilities of the variants are reported in Fig. 1. Kinetic constants ($K_m$, $k_{cat}$) for propane oxidation were determined from Michaelis–Menten plots of initial

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**Table 1. Summary of directed evolution steps and amino acid substitutions accumulated in P450BM3 variants**

<table>
<thead>
<tr>
<th>Variant</th>
<th>Target substrate</th>
<th>Mutagenesis method</th>
<th>Heme domain substitutionsa</th>
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<td>1-3e</td>
<td>Propane</td>
<td>Site saturation—recombination</td>
<td>V138H, I175V, F205C, S226R, T295A</td>
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<tr>
<td>Jc</td>
<td>Propane DNA shuffling</td>
<td>Site directed</td>
<td>V178I, A184V, H236Q, A245S, A263S, T295A</td>
</tr>
<tr>
<td>19A12a</td>
<td>Propane</td>
<td>Random</td>
<td>L52I, I366V</td>
</tr>
<tr>
<td>11-3f</td>
<td>Propane Site saturation</td>
<td>Recombination</td>
<td>L188P</td>
</tr>
<tr>
<td>P450PMO</td>
<td>Propane</td>
<td>Saturation—recombination/site directed</td>
<td>A328F</td>
</tr>
</tbody>
</table>

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*Substitutions are given with respect to the preceding variant.

a From Ref. 32.

b From Ref. 27.

c From Ref. 28.

d From Ref. 30.

† From Ref. 27.

From Ref. 29.

†† From Ref. 31. This and subsequent variants carry also two mutations in the reductase domain (E464G, I710T). While important for ethane oxidation, these mutations have only a small effect on propane oxidation.

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† † http://drnelson.utmem.edu/P450.statsfile.html
propanol formation rates at different propane concentrations. Since saturation of 139-3 and J could not be achieved within the limits of propane solubility in water (≈1.6 mM), the kinetic parameters determined for these variants are approximate. Compared to 139-3 (wild-type P450BM3 has no detectable propane activity), total turnovers supported by P450PMO increased 170-fold (Fig. 1a). This increase was accompanied by a 180-fold decrease in $K_m$ (from ~30 mM in 139-3 to 170 μM in P450PMO; Fig. 1b) and by a 45-fold increase in $k_{cat}$ (from ~10 min$^{-1}$ in 139-3 to 450 min$^{-1}$ in P450PMO; Fig. 1c). Overall, P450PMO shows a catalytic efficiency ($k_{cat}/K_m$) on propane of 4.4×10$^4$ M$^{-1}$ s$^{-1}$, which is approximately 4 orders of magnitude greater than that of the first propane oxidizer, 139-3 (~5 M$^{-1}$ s$^{-1}$).

Despite the much reduced size and apolar nature of propane, P450PMO’s $K_m$ for propane oxidation is comparable to that of P450BM3 for laurate oxidation (130 μM), while its catalytic efficiency is only four times lower (cf. 1.9×10$^5$ M$^{-1}$ s$^{-1}$).

The coupling efficiency for propane oxidation improved from only 5–8% in the earliest generations to 93% in P450PMO (Fig. 1d). The largest change in coupling efficiency (15% → 44%) was caused by the single L188P mutation between ETS8 and 19A12. Coupling of product formation to NADPH oxidation approaches 100% only with ideal substrates for which no competitive side reactions occur during the catalytic cycle. Thus, the increasing coupling efficiency along the variant series suggests that the active site is becoming more optimized for propane binding. This conclusion is supported by the kinetic data and is consistent with the large decrease in $K_m$.

How the evolution of propane activity affects the native activity was evaluated by measuring variants’ activity on the long-chain fatty acids palmitate (C16) and laurate (C12) and comparing those to that of wild-type P450BM3 (Fig. 1e). P450PMO has no detectable activity on laurate and only marginal activity on palmitate ($b_150$ TTN, or less than 0.5% that of P450BM3). Palmitate is a significantly better substrate for P450BM3 than laurate in terms of catalytic efficiency (6.0×10$^7$ M$^{-1}$ s$^{-1}$ versus 1.9×10$^5$ M$^{-1}$ s$^{-1}$), which probably explains why laurate activity was lost completely during P450PMO evolution. As Fig. 1e shows, native function was only mildly affected in the first two intermediates, 139-3 and J [J displays 31% (palmitate) and 61% (laurate) activity relative to wild type]. A more significant decrease was observed with 35E11 (~10% relative activity).

The mutations accumulated in 35E11 (~10% relative activity). The mutations accumulated in 35E11 include R47C (Table 1), which removed a side chain important for recognition of the carboxylate moiety in fatty acid substrates. We reverted Cys47 to Arg in both 35E11 and P450PMO to examine its contribution to the loss of fatty acid activity. In 35E11, C47R led to moderate recovery of activity (laurate, 8–24% relative activity; palmitate, 5–18% relative activity), whereas P450PMO-C47R still had no measurable activity on laurate and negligible activity on palmitate (~180 TTN). These results indicate that refinement of propane activity came at the cost of the

Fig. 1. Functional and kinetic properties of P450PMO evolutionary intermediates. (a) Total turnover numbers for propane hydroxylation (moles propanol per mole P450). (b) $K_m$ values and (c) catalytic rate constants ($k_{cat}$) for propane oxidation. (d) Coupling efficiency as given by propanol formation rate/NADPH oxidation rate. Errors are within 10%. (e) Relative activity with palmitate and laurate as compared to P450BM3. (f) Enzyme thermostability, reported as $T_{50}$ values from heat inactivation of heme domain after 10 min incubation. (g) Average stability penalty per amino acid substitution for each step in the lineage.
original function regardless of the presence of specific fatty-acid-binding determinants. In our experiments, no detectable activity on propane or fatty acids corresponds to a catalytic efficiency of less than 1 M$^{-1}$ s$^{-1}$. Therefore, from P450$^{\text{BM3}}$ to P450$^{\text{PMO}}$ the overall change in specificity is at least a factor of $(4.4 \times 10^{4}/1)/(1/1.9 \times 10^{5}) \sim 10^{10}$.

To establish the impact of the accumulated mutations on enzyme stability, $T_{50}$ values were determined from heat-inactivation curves of the heme-dependent CO-binding spectral shift (Fig. 1f). These analyses show a gradual but continual decrease in the heme domain stability along the evolutionary pathway, reflecting the fact that stability was not under selection. The average impact of each mutational event on stability varies considerably though, as shown by the plot of per-mutation stability penalties (Fig. 1g). With 35E11, the much reduced stability compared to wild type ($\Delta T_{50}=−11.6$ °C) prevented further evolution and necessitated the stabilization step that led to ETS8, at which point a new beneficial mutation (L188P) could be found. To investigate whether L188P exerts the same effect in the background of the less stable precursor, we introduced L188 in 35E11, where it caused a perturbation in the heme environment as indicated by the appearance of a 420-nm peak (~40%) in the CO-difference spectrum of the purified enzyme (data not shown). Furthermore, in contrast to the ~300% increase in TTN$_{\text{propane}}$ as a result of Leu188 mutation to proline during ETS8→19A12 transition, L188P generated only a small improvement (~40%) in 35E11. Altogether, these results highlight the role of the stabilizing mutations in enabling the fold to accommodate the beneficial but destabilizing L188P substitution. Saturation mutagenesis of position 188 in 19A12 confirmed proline as the best possible amino acid substitution (data not shown).

**Substrate specificity profiles**

Specificities of the evolved P450 variants were investigated using the alkane series from C$_{3}$ (ethane) to C$_{10}$ (decane). Total turnovers—a cumulative measure of both catalytic and coupling efficiency—were determined for each substrate→P450 pair under identical reaction conditions (Supplementary Table S1). To enable comparison, TTNs were normalized to the highest value in the series (maximal turnover number). The resulting profiles (Fig. 2a) reveal a remarkable shift in substrate preference along the evolutionary path. The activity of P450$^{\text{BM3}}$ on the C$_{2}$–C$_{10}$ alkanes was marginal (maximal turnover number=160) and restricted to C$_{8}$ > 8 alkanes. In the earliest steps, substrate preference centered on the medium-chain alkanes (C$_{8}$–C$_{9}$), extending only later (35E11) toward shorter alkanes. A change in substrate preference appeared with 19A12. Subsequent variants 11-3, 1-3, and P450$^{\text{PMO}}$ showed a progressive decrease in TTN with alkane length. In P450$^{\text{PMO}}$, specificity along the alkane series has become remarkably restricted around propane: TTN drops by 55% for butane and by 95% for pentane.

Initial rates of product formation were compared for C$_{3}$ to C$_{6}$ alkanes (Fig. 2b). In the earliest intermediates, the preference for longer-chain alkanes is indicated by high $v_{\text{a}}$(C$_{9}$)/$v_{\text{a}}$(C$_{3}$) ratios (51 for 139-3; 14 for J). This ratio decreased to ~1 in 35E11 and then switched in favor of the smaller alkane in subsequent generations. In P450$^{\text{PMO}}$, the product formation rate was highest on propane and decreased along the alkane series ($v_{\text{a}}$(C$_{9}$)/$v_{\text{a}}$(C$_{3}$)=0.039). Overall, the ratio of the rates on propane versus nonane changed 1300-fold from 139-3 to P450$^{\text{PMO}}$.

Upon substrate binding and displacement of the water ligand from the heme iron in the low-spin configuration, ferric heme converts to a high-spin configuration, which appears as a blue shift of the Soret band from ~41 to ~392 nm. How the variants interact with alkanes was further investigated by testing the heme iron spin shifts in the presence of propane, pentane, and octane (Fig. 2c). High-spin content was estimated based on reference spectra of P450$^{\text{BM3}}$ in complex with increasing amounts of palmitate. Octane binding induced between 35% and 50% spin shift in 139-3 and later-generation variants. Significant propane- and pentane-induced spin shifts (~25%) were observed only in the last three variants studied. Interestingly, 19A12 and its successors showed partial high-spin content (5–10%) even in the absence of substrate, suggesting that the mutations displace water in the active site or interfere with hydrogen bonding to the axial water ligand.

In a further probe of the active site, we also tested P450$^{\text{PMO}}$ and its precursors on three structurally diverse terpenes: guaiol, valencene, and limonene. Oxidation products were quantified by GC (Fig. 3). For all variants, the TTN changed inversely with substrate size (guaiol ≪ valencene ≪ limonene). Early intermediates 139-3 and particularly J showed the highest activities with all three, demonstrating substrate promiscuity that extends beyond alkanes. A large drop in activity was observed in the J→35E11 transition, followed by a gradual but steady decrease as activity on propane increased. P450$^{\text{PMO}}$ behaved similarly to P450$^{\text{BM3}}$, showing negligible activity on guaiol and valencene (TTN ≪150) and very weak activity on limonene (TTN ≈580).

**Halomethane dehalogenase activity**

P450$^{\text{PMO}}$ supports thousands of turnovers on ethane (TTN ~2450) despite the higher C–H bond strength (101.0 kcal mol$^{-1}$ for ethane versus 98.6 kcal mol$^{-1}$for the subterminal C–H of propane). To determine whether P450$^{\text{PMO}}$ can insert oxygen into C–H bonds stronger than those of ethane, P450$^{\text{PMO}}$ activity was tested on halomethanes (CH$_{3}$Cl, CH$_{3}$Br, and CH$_{3}$I), which have molecular sizes and C–H bond energies intermediate between those of methane and propane. These reactions with halomethanes were quantified by GC (Supplementary Fig. S2).
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Fig. 2. Evolution of a P450 propane monoxygenase from a long-chain fatty acid hydroxylase. (a) Substrate specificity profiles of P450<sub>PMO</sub> and its evolutionary intermediates on the C<sub>2</sub>–C<sub>10</sub> alkane series. Maximal turnover numbers: P450<sub>BM3</sub> = 160; 139-3 = 2100; J = 3450; 35E11 = 6350; 19A12 = 10,500; 11-3 = 13,200; 1-3 = 19,200; P450<sub>PMO</sub> = 33,400. The dashed line indicates the profile of the preceding variant. Numbers indicate the number of heme domain mutations with respect to P450<sub>BM3</sub> and the preceding variant (brackets). The chemical structures next to the arrows refer to the selective pressure (octane or propane). The preferred substrates of P450<sub>BM3</sub> (palmitate) and P450<sub>PMO</sub> (propane) are displayed as space-filling models. (b) Initial oxidation rates with different alkanes, as measured by total nanomoles of hydroxylated alkane per nanomole enzyme per minute. (c) Alkane-induced shifts of heme spin state with different substrates, as determined from 350 to 500 nm spectra.

Fig. 3. Activities of P450<sub>PMO</sub> evolutionary intermediates on terpene substrates, as determined by GC analysis of the oxidation products.
expected α-elimination of HX upon hydroxylation. Measurable bromomethane activity first appeared in 35E11 and increased through the series at a level that is 1–5% of the propane activity (Fig. 4b). Despite the \( \sim 1 \text{ kcal mol}^{-1} \) weaker C–H bond, chloromethane activity emerged only later (19A12) and was detected at lower levels (0.5–1% of propane activity), reflecting the preference of these variants for propane-sized substrates. Unlike CH₃Cl and CH₃Br, iodomethane is not oxidized by mammalian P450s.⁴⁰ Interestingly, P450PMO and its precursor 1–3 exhibited detectable activity on iodomethane, albeit with low turnover rates (0.5–1 min⁻¹). These results provide the first demonstration that a 103.0 kcal mol⁻¹ sp³ C–H bond is energetically accessible to oxygen insertion by a P450 enzyme.

Structural investigation of the P450PMO lineage

Crystallization experiments were undertaken to investigate structural changes along the lineage. While P450PMO and other variants failed to yield diffraction-quality crystals, early intermediate 139-3 could be crystallized in the presence of N-palmitoylglycine (NPG). The structure of the complex was solved at 2.6 Å resolution with \( R = 0.21 \) and \( R_{	ext{free}} = 0.26 \) (see Table 2 for data collection and refinement statistics). The asymmetric unit contains two chains, A and B. The structure of P450bm3 in complex with NPG is available.⁴¹ Superposition of the 139-3 and P450bm3 complexes revealed no gross changes in the enzyme fold (Fig. 5a, average Cα RMSD = 0.5 Å). The largest deviations (>1 Å) were in the N-terminal region (Thr1–Leu20), the loop in β-sheet β1 (Pro45–Arg47), the region between helix H and I (Pro243–Glu247), and the 3₁₀ helix between Glu380 and Ser383 (Fig. 5a and Supplementary Fig. S3). The NPG ligand was found to adopt a more extended conformation in complex with 139-3, and a slight movement of β-sheet β1 toward helix F caused a restriction at the entrance of the substrate access channel (Fig. 5a). The relative positions of the residues critical for fatty acid interaction (Arg47, Tyr51) were unaltered. V78A and A184V are in the active site and are at short distance from one another (Fig. 5b). V78A is responsible for expansion of the hydrophobic pocket near the heme, whose volume increased by 29 Å³ (Fig. 6a). In 139-3, Val1184 forms a much tighter (3.3 Å versus 4.2 Å) contact with Leu437 across the substrate channel. Assuming octane orients similarly to NPG in the active site, Leu437 would establish closer van der Waals contacts near the terminal carbon of octane and enable more efficient exclusion of the shorter substrate from the solvent. This also translates into a change of the substrate orientation in the active site as reflected by the regioselectivity for octane oxidation in 139-3, which hydroxylates primarily at the 2-position (55% 2-octanol, 20% 3-octanol, 18% 4-octanol, and 7% ketone) versus the

Table 2. Data collection and refinement statistics

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<td>Total observations</td>
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Backbone geometry analysis revealed only one residue (Leu44³) in a disallowed region of Ramachandran plot, which is nevertheless very well defined with clear density.

Footnotes:

⁶ Values in parentheses are for the highest-resolution shell.
⁷ \( R_{sym} = \Sigma |I|/\Sigma l \) / \( \Sigma |I| \), where \( I \) is the observed intensity and \( l \) the averaged intensity of multiple symmetry-related observations of the reflection.
⁸ \( R_{free} = \Sigma |I|/\Sigma l \) / \( \Sigma |F_o| \), where \( F_o \) and \( F_e \) are the observed and calculated structure factors, respectively.
⁹ \( R_{free} \) was calculated with the 5% of reflections set aside randomly throughout the refinement.
wild-type enzyme, which prefers the 3- and 4-positions (5% 2-octanol, 49% 3-octanol, 43% 4-octanol, and 3% ketone).

Conservative structural models of 35E11, 19A12, 1-3, and P450PMO were calculated based on the 139-3 crystallographic model (chain B) using the Rosetta structure prediction algorithm. In 35E11, three mutations (V78F, A82S, A328F) cause a major change in the substrate channel. Specifically, Phe328 (in concert with Leu437 and Ala74) constricts the

Fig. 5. Structure of variant 139-3. (a) Superposition of NPG-bound structures of P450BM3 (orange, PDB code 1JPZ) and evolutionary intermediate 139-3 (green, this work; PDB code 3CBD). The mutated residues, fatty acid-binding Arg47 and Tyr51, and Leu188 are displayed as stick models. The heme is displayed as a space-filling model. N- and C-termini and selected secondary-structure elements are indicated with capital letters. (b) Contacts of V78A and A184V substitutions with Leu347 and the bound ligand in 139-3 (green). (c) Superposition of 139-3 structure (green) and 19A12 model (red) illustrating the helix-capping role of Leu188 and the disruption of helical conformation as a consequence of the L188P mutation in 19A12.

Fig. 6. Substrate access pathway. (a) Volumetric restriction and compartmentalization of the substrate access pathway during the transition of P450BM3 to P450PMO (left). The A328F mutation in 35E11 interrupts the contiguity of the path by defining an inner cavity (at the heme pocket) from the remaining of the access path (indicated as ‘substrate channel’). The computed fraction of polar surface area (right) shows an increasing difference in polarity between the inner and outer compartments along the lineage. (b) Surface representations of the substrate access pathway in P450BM3 structure, 139-3 structure, and P450PMO structural model. The cavity above the heme first expands to accommodate octane (in 139-3). Upon selection for propane activity, this space becomes narrowly confined by a phenylalanine triad (black arrows) and segregated from the outer substrate channel through the interaction of Leu347 with opposing Glu74.
channel so that the contiguous substrate access pathway found in P450BM3 and 139-3 (Fig. 6b) appears compartmentalized into an inner heme pocket (HP) and a more solvent-accessible substrate channel (SC) (as mapped using a probe radius of 1.4 Å). The total volume of these cavities is dramatically reduced compared to that of 139-3 ($\Delta V_{\text{SC + HP}} = -127$ Å$^3$; Fig. 6a). Experimental data support these calculations. The $>$70% drop in activity toward the terpenes suggests a much reduced accessibility of 35E11’s active site compared to those of the preceding variants (139-3, J), while the increased regioselectivity with C$_7$–C$_8$ alkanes ($>$90% 2-alkanol) is indicative of a significant decrease in these substrates’ conformational freedom within the active site.

In contrast to the other mutations, L188P was associated with a very unfavorable energy. The structure of 19A12 was therefore calculated allowing backbone motion for residues 185 to 191 and updating the conformation of this region using Loopy. In the resulting model, Pro188 causes partial unwinding of helix F C-terminus and removes Leu188 steric bulk, reducing the distance between helix F and adjacent helix B’ (Fig. 5c). These changes appear to have significant impact on the properties of the enzyme (see Discussion). In P450$_{\text{PMO}}$, the active-site mutations A184V, S74E, and S82G further restrict the substrate channel (Fig. 6a). This enables a more productive interaction of the enzyme with the substrate as indicated by the higher high-spin content upon propane binding and increased coupling in propane oxidation. The G443A mutation was found to affect the propane oxidation rate (twofold) but has little effect on coupling (91%) and $K_m$ for propane (170 μM).

**Discussion**

Propane is one-fifth the size of a typical cytochrome P450$_{\text{BM3}}$ substrate and lacks the carboxylate moiety and long aliphatic chain through which the fatty acid interacts with the enzyme (Fig. 2a). By studying the intermediates along the pathway to P450$_{\text{PMO}}$, we were able to observe how the enzyme has reelaborated its substrate-recognition strategy to achieve catalytic proficiency on this gaseous alkane. The 139-3 crystal structure shows that broadening of the substrate range of P450$_{\text{BM3}}$ to include octane and other medium-chain alkanes occurred with minimal perturbation of the overall fold. An enlarged hydrophobic pocket near the heme (Fig. 6a and b) appears to provide an energetically more favorable environment for binding to octane while also enabling the emergence of propane activity. These changes interfered only slightly with the native enzyme activity, which decreased by a factor of ~2. The latter reflects a previously observed feature of enzyme evolution: improvement of side, or ‘promiscuous,’ activities under positive selection pressure often comes with only marginal cost to the native activity. The substrate range of variants 139-3 and J appears to be broad and includes other hydrophobic substrates (i.e., terpenes) that differ considerably from those used for activity screening (octane and propane, respectively). This generalist character has been observed in other early products of directed evolution such as, for example, in engineered variants of a beta-glucuronidase.

In the next few generations, the prevailing strategy for improvement of propane activity involved compartmentalization and restriction of the active-site cavity (cf. 35E11). This facilitated the interaction of the enzyme with the smaller substrate, as evidenced by (a) a sevenfold increase in TTN$_{\text{propane}}$, (b) appearance of the heme spin shift (8%) upon propane binding, and (c) a decrease in $K_m$ to the submillimolar range. The substrate preference remained centered on C$_7$–C$_8$ alkanes, but a significant decrease ($>$90%) in the native function was observed. The ‘turning point’ in substrate specificity in the next step (35E11→19A12) was induced by a single amino acid substitution, L188P (ETS8’s profile overlaps with that of 35E11; data not shown). In the 139-3 structure, Leu188 lies at the C-terminus of helix F that, together with helix G, forms a lid that undergoes conformational change upon substrate binding (Fig. 5a). L188P removes the intrastrand H bonds established by the Leu188 amide NH and is associated with a perturbation of the surrounding helical region (Fig. 5c) and a large destabilizing effect ($\Delta T_{50} = -3.3$ °C versus mean value of $-0.6$ °C/mutation, Fig. 1g). The stability cost of this mutation is offset by the advantage—the largest on a per-residue basis—it provides in propane oxidation, reflected in a threelfold improvement in TTN and coupling efficiency, a twofold increase in turnover rate constant $k_{\text{cat}}$, and a twofold reduction in $K_m$ (830 μM in ETS8 versus 475 μM in 19A12). By bringing helix F and helix B’ in closer proximity, this mutation may force the enzyme to adopt a conformation similar to the closed, catalytically more favorable one.

After 19A12, a gradual decrease in activity towards longer alkanes accompanied the increased activity on propane, even though no negative selection was applied during the evolution of these variants to specifically favor this trend. Mapping of the substrate access pathway revealed a twisted path connecting the exterior to the heme (Fig. 6b). This configuration apparently provides an effective solution to channeling propane to the active site. It is also likely to hamper diffusion of longer-chain alkanes to and from the active site, explaining the observed difference in oxidation rates and TTNs observed within the alkane series. A comparison can be made with the soluble non-heme diiron methane monoxygenase (sMMO), an evolutionarily unrelated enzyme found in methanotrophic bacteria. Studies addressed the question of how sMMO can discriminate methane from hundreds of other potential substrates with weaker C–H bonds.

The α subunit of sMMO hydroxylase domain was found to direct methane to the active-site pocket through a sequence of five buried cavities.
remarkable restriction ($\Delta V = -248 \text{ Å}^3$; Fig. 6a) and compartmentalization of the substrate access pathway along the P450PMO lineage suggest a similar adaptive strategy to dealing with the problem of recognizing a small gaseous substrate. This can be rationalized in light of the negligible binding energy contribution ($-\Delta G < -4 \text{ kcal/mol}$) provided by neutral ligands with fewer than five non-hydrogen atoms and the fact that propane presumably has insufficient binding energy to induce the lid closure and accompanying conformational changes that characterize the interaction of P450BM3 with long-chain fatty acids. In contrast to the highly hydrophobic 'chambers' in sMMO, however, the substrate access channel in P450PMO has a more pronounced polar character. In particular, a progressive partitioning between a hydrophobic heme pocket and a relatively more polar substrate channel was observed along the variant series (Fig. 6a), a trend that culminated with a charged residue (Glu74) interposing between the two (Fig. 6b). Noteworthy in this regard was the progressive increase in polarity and bulkiness of residue 74 over the final evolutionary steps: Ala (19A12) $\rightarrow$ Ser (1-3) $\rightarrow$ Glu (P450PMO).

While optimal for oxidation of the gaseous substrate, the acquired changes are incompatible with the native and other activities. In P450PMO, there remains little trace of the original P450BM3 function. Furthermore, in contrast to the earliest intermediates, which exhibit broad substrate ranges, P450PMO appears to have completely respecialized, displaying very narrow substrate tolerance within the C$_2$-C$_{10}$ alkane series and little activity toward other, unrelated substrates (Figs. 2 and 3). Thus, the respecialization of P450PMO is induced by the same forces that pushed for optimization of the function under selective pressure and appears to be a consequence of acquisition of catalytic proficiency with propane.

Altogether, our results have several important implications. First, these data show that far from being 'evolutionary dead ends,' specialized enzymes can serve as starting points for divergent trajectories that lead to other, specialized members of the same enzyme family. Also, we provide the first experimental evidence that positive selection alone is sufficient to induce complete respecialization of a P450 enzyme, which suggests that negative selection need not have played an important role in the evolution of specialized members of this and possibly other enzyme families. Finally, the emergence of iodo- methane dehalogenase activity in P450PMO is noteworthy, as mammalian P450s and sMMO do not exhibit it. Iodomethane is known to be degraded by a very small subset of methylotrophic species, using enzymes such as corrinoid-dependent methyltransferases. The propane-adapted features of P450PMO's active site enable the emergence of a basal activity towards this substrate, thus providing an alternative enzymatic strategy for the degradation of this compound and further expanding the range of known P450 activities.

Materials and Methods

Kinetic, biochemical, and biophysical analyses

Enzymes were expressed and purified as previously described. P450PMO, P450BM3(C47R), 35E11(C47R), and 35E11(L188P) were obtained by site-directed mutagenesis (see Supplementary Table S2 for primer sequences). Michaelis–Menten plots for propane oxidation (e.g., Supplementary Fig. S1) were obtained varying propane concentration by dilution of propane-saturated solutions ($\approx 1.6 \text{ mM}$). Reactions (5 mL) were carried out at 25 °C in 100 mM KPi buffer (pH 8.0) using 100–400 nM enzyme and 1 mM NADPH. After 20 s, reactions were stopped with 200 μL concentrated H$_2$SO$_4$ and analyzed by gas chromatography–electron capture detector (GC–ECD) as previously described. Initial rates (20 s) for alkane oxidation were measured using 1 mM substrate (2% ethanol), 50 nM P450, and 1 mM NADPH. Reactions were stopped with 100 μL 2 N HCl, extracted with CH$_3$Cl$_2$, and analyzed by GC–flame ionization detector (GC–FID). Coupling efficiency corresponds to the ratio between propanol formation rate and NADPH oxidation rate in the presence of propane and was measured as previously described. Half-denaturation temperatures ($T_{sd}$) were determined as follows. Samples of purified enzyme (~3 μM) were incubated for 10 min at different temperatures (from 20 to 60 °C) in a PCR thermocycler. After centrifugation, 160-μL protein solutions were mixed with 40 μL 0.1 M sodium hydrosulfite on a microtitre plate and incubated for 10 min with carbon monoxide in a sealed chamber. $T_{sd}$ values were calculated from heat-inactivation curves of CO-binding difference spectra. Alkane-induced spin shifts were recorded on a Shimadzu UV–VIS spectrophotometer using P450 (2 μM) and 1 mM octane, pentane, or propane (2% ethanol). High-spin content was estimated by comparison with spectra of P450BM3 and palmitate mixtures. Reactions with alkanes from pentane to decane were carried out at 3-mL scale using KPi buffer containing 2% ethanol, 50 nM enzyme, 1.6 mM alkane, and a cofactor regeneration system (CRS: 10 U/mL isocitrate dehydrogenase, 7 mg/mL isocitrate, 0.15 mg/mL NADP$^+$). Reaction mixtures were sealed, stirred for 24 h (4 °C), extracted with CH$_3$Cl$_2$ and analyzed by GC–FID. Reactions with ethane, propane, and butane were carried out as described for C$_5$–C$_{10}$ alkanes using alkane-saturated buffers. All kinetic and biochemical measurements were performed in triplicate. For C$_5$–C$_{10}$ alkane reactions, GC analyses were carried out using a Shimadzu GC-17A gas chromatograph, an Agilent HP5 column (30 m $\times$ 0.32 mm $\times$ 0.1 μm film), 1 μL injection, FID detector, and the following program: 250 °C inlet, 50 °C (3 min), 5 °C/min to 100 °C, 25 °C/min to 230 °C, and 230 °C (3 min). For butane and propane reactions (TTN), GC analyses were carried out using a Hewlett-Packard 5890 Series II Plus gas chromatograph, a Supelco SPB-1 column (60 m $\times$ 0.32 mm $\times$ 0.1 μm film), 0.5 μL injection, FID detector, and the following program: 250 °C inlet, 80 °C (2 min), 10 °C/min to 110 °C, 25 °C/min to 275 °C, and 275 °C (2.5 min).

Analysis of fatty acid and terpene activity

Reactions with laurate and palmitate were carried out at 2-mL scale using 2 mM fatty acid, 100–500 nM enzyme, and CRS. After stirring for 24 h, solutions were acidic with HCl, extracted with hexane, and dried over Na$_2$SO$_4$. N-Methyl-N-trimethylsilylheptafluorobutylamide 1:10
After 30 min reaction, 1-mL aliquots were mixed with 500 nM enzyme, and CRS. Buffers were prepared with NaHPO4 (pH 7.4), 100 mM NaCl, 10 mM β-mercaptoethanol (buffer A). All activity measurements were performed in triplicate.

Reactions with halomethanes

Reactions were carried out at 1-mL scale using halomethane-saturated 100 mM KPi (pH 8.0), 50 to 500 nM enzyme, and CRS. Buffers were prepared with HPLC-grade water and autoclaved twice prior to use. After 30 min reaction, 1-mL aliquots were mixed with 50 μL o-(2,3,4,5,6-pentafluorobenzyl)-hydroxylamine hydrochloride (Sigma) in formaldehyde-free water (17 mg/mL) and incubated for 15 min at 25 °C. Mixtures were mixed with 10 μL 25 mM 1-bromohexane (internal standard) and extracted with 500 μL hexane. The organic phase was diluted 1/100 (v/v) and analyzed by GC–FID using a Shimadzu GC-17A gas chromatograph, an Agilent HP5 column, 1 μL injection, and the following program: 300 °C inlet, 100 °C (2 min), 15 °C/min to 200 °C, 50 °C/min to 220 °C (3 min). A calibration curve was constructed from 2 μL injection, FID detector, and the following program: 70 °C inlet, 45 °C (1 min), 15 °C/min to 200 °C, 50 °C/min to 220 °C, and 220 °C (5 min). A calibration curve was constructed from freshly prepared solutions of formaldehyde. All measurements were performed in triplicate.

Protein crystallization

For crystallization, the heme domain of 139-3 (1–455 residues; 139-3H) was cloned into the EcoRI/KpnI cassette of vector pProEX-1 using 5′-GATCCGGAATTCATGA-3′ and 5′-GCTAAAAAGTA-TAGGTACCAAGCCTTGCC-3′ as forward and reverse primers, respectively. Recombinant Escherichia coli DH5α cells were grown in terrific broth medium and induced with 0.6 mM IPTG at OD600 0.7. Flasks were transferred to 25 °C and harvested after 20 h. Cells were lysed in 50 mM sodium phosphate (pH 7.4), 100 mM NaCl, 1 mM PMSE, and 10 mM β-mercaptoethanol (buffer A). Cell lysate was loaded onto a Ni–NTA agarose column preequilibrated with buffer A. After the column was washed with 50 mM NaHPO4 (pH 7.4), 100 mM NaCl, 10 mM β-mercaptoethanol, and 10% glycerol (buffer B), bound protein was eluted with buffer B in the presence of 100 mM imidazole. After concentration, the protein was loaded onto a Sephacryl S200 gel-filtration column reequilibrated with buffer A. Purified 139-3H was mixed with NPG and crystallized by hanging-drop vapor-diffusion method from a protein solution at 40 mg/mL and a solution containing 100 mM Mops (pH 7.0), 50 mM LiBr, and 20% PEG (polyethylene glycol) 20,000. Crystals were soaked in 2 M sodium malonate as cryoprotectant prior to data collection.

Data collection and structure determination

X-ray diffraction data were collected on beam line 5.0.1 at Advanced Light Source (Berkeley, CA) using an ADSC Quantum 4 CCD detector. Optimization of data collection was guided by the STRATEGY function of MOSFLM. All data were reduced using DENZO and scaling and rejections were performed with SCALEPACK. The structure was solved using molecular replacement as implemented in AMORE in CCP4 suite and the structure of P450BM3 complexed with palmitoleic acid (PDB ID code 1FAG) as search model. Two solutions in the asymmetric unit were found using data between 43.0 and 3.0 Å. Phases were improved and extended to 2.65 Å over 50 iterations of density modification using DM as well as twofold noncrystallographic symmetry averaging. The structure was further refined using the graphical model building program O and CNS to a final resolution of 2.65 Å. Backbone geometry was checked in PROCHECK. Data collection and refinement statistics are provided in Table 2.

Model calculations and substrate access pathway analysis

Structural models were calculated with Rosetta using the structure of intermediate 139-3 (chain B) as a reference model. The positions of mutated side chains were optimized (including off-rotamer continuous optimization) using full-atom Rosetta energy function. To accommodate L188P, the backbone between residues 185 and 191 was updated using Loopy. Substrate access pathways were analyzed with Quick hull using Cys400 and C atoms of residues 72–75, 78, 82, 87, 88, 177, 182, 185, 259, 263–268, 328–332, 357, and 435–439 as reference points and constructing a convex hull around these points. Within the hull, a ‘castr’ was formed by placing dummy atoms at all 0.5 Å grid points at least 3 Å from a protein atom. Substrate pathways were visualized displaying the surface of the cast with PyMOL. Solvent-excluded volumes were computed using MSMS with a probe radius of 1.4 Å and a density of 3.0/Å2. Reported volume and surface area values refer to volume occluded and the area buried, respectively. Based on MSMS per atom surface area breakdown, polar and nonpolar surface areas were measured using OPLSa charges and grouped atoms with a weight <0.2 into the nonpolar category.

Protein Data Bank accession number

Coordinates and structure factors have been deposited in the Protein Data Bank with accession number 3CBD.

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† www.pdb.org
§ http://www.pymol.org
Supplementary Data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2008.06.060

References


