*Pseudomonas aeruginosa* cytochrome P450 CYP168A1 is a fatty acid hydroxylase that metabolizes arachidonic acid to the vasodilator 19-HETE

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*Pseudomonas aeruginosa* is a Gram-negative opportunistic human pathogen that is highly prevalent in individuals with cystic fibrosis (CF). A major problem in treating CF patients infected with *P. aeruginosa* is the development of antibiotic resistance. Therefore, the identification of novel *P. aeruginosa* antibiotic drug targets is of the utmost urgency. The genome of *P. aeruginosa* contains four putative cytochrome P450 enzymes (CYPs) of unknown function that have never before been characterized. Analogous to some of the CYPs from *Mycobacterium tuberculosis*, these *P. aeruginosa* CYPs may be important for growth and colonization of CF patients’ lungs. In this study, we cloned, expressed, and characterized CYP168A1 from *P. aeruginosa* and identified it as a subterminal fatty acid hydroxylase. Spectral binding data and computational modeling of substrates and inhibitors suggest that CYP168A1 has a large, expansive active site and preferentially binds long chain fatty acids and large hydrophobic inhibitors. Furthermore, metabolic experiments confirm that the enzyme is capable of hydroxylating arachidonic acid, an important inflammatory signaling molecule present in abundance in the CF lung, to 19-hydroxyeicosatetraenoic acid (19-HETE; *K*<sub>m</sub> = 41 μM, *V*<sub>max</sub> = 220 pmol/min/nmol P450), a potent vasodilator, which may play a role in the pathogen’s ability to colonize the lung. Additionally, we found that the *in vitro* metabolism of arachidonic acid is subject to substrate inhibition and is also inhibited by the presence of the antifungal agent ketoconazole. This study identifies a new metabolic pathway in this important human pathogen that may be of utility in treating *P. aeruginosa* infections.

*Pseudomonas aeruginosa* is a Gram-negative opportunistic pathogen that is highly prevalent in individuals with cystic fibrosis (CF), a debilitating inherited disease (1–3). In CF patients with reoccurring *P. aeruginosa* infections, the organism is known to form intractable biofilms that lead to antibiotic resistance (4–7). Antibiotic failure can result in pneumonia, which can be life-threatening in the respiratory compromised CF patient (2, 8). Moreover, chronic airway infection by *P. aeruginosa* significantly promotes lung tissue destruction, further compromising pulmonary function in individuals with CF. Given this, the need for characterization of new antibiotic targets and drugs that have the potential to inhibit biofilm formation in *P. aeruginosa* is urgently needed. The genome of *P. aeruginosa* strain PAO1 (UW) was first completely sequenced in 1999 (9) and serves as a reference genome for the organism (http://www.pseudomonas.com/). The genome of *P. aeruginosa* contains four putative cytochrome P450 (CYP) monooxygenase genes: designated CYP107S1, CYP168A1, CYP169A1, and CYP239A1 (10). In bacteria, CYP enzymes are known to perform diverse functions, including: antibiotic synthesis (11–14), carbon source metabolism (15–17), detoxification (18, 19), and secondary metabolite production (20–22). CYP enzymes from the closely related obligate intracellular pathogen *Mycobacterium tuberculosis* have been characterized and determined to be integral to the metabolism of fatty acids (23–26), cholesterol (27, 28) and even certain antibiotics (29, 30), all of which promote bacterial survival and growth under a variety of conditions. While the exact functions of the CYP enzymes identified in *P. aeruginosa* remain unknown, they have recently been implicated in both detoxification (18) and the oxidation of environmental contaminants (31). Further evidence links them to the metabolism of medium- to long-chain alkanes (32), suggesting their role as fatty acid and/or alkane hydroxylases. Indeed, a number of bacterial CYP enzymes are known to function as fatty acid hydroxylases, and this role for CYPs seems to be common among the eubacteria (10), although this particular function has yet to be demonstrated for the *P. aeruginosa* CYP enzymes.

Many types of fatty acids are found within the environment of the lung and could possibly serve as carbon sources for *P. aeruginosa* as it attempts to establish a colony (8). However, certain fatty acids, including arachidonic acid and its metabolites, control inflammation and immune cell recruitment to the site of infection (33–35). Moreover, concentrations of arachidonic acid are significantly increased in the CF patient lung due to a metabolic defect (36, 37). An intriguing possibility is that the organism may be able to modify the immune response by oxidation of arachidonic acid and its metabolites, thereby making its local environment more hospitable for the parasite to grow and proliferate. Indeed, a secreted
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lipoxygenase from P. aeruginosa, designated LoxA, is capable of metabolizing arachidonic acid to 15-hydroxyeicosatetraenoic acid (15-HETE) (38), which has been demonstrated to downregulate the immune response in vivo (39), illustrating the capacity of P. aeruginosa to reduce the host's ability to respond to infection through the metabolism of inflammatory mediators. More recent work elucidated the role of a soluble epoxide hydrolase to specifically reduce levels of the host proresolving lipid mediator, 15-epi lipoxin A4 (15-epi LXA4), thereby contributing to the prolongation of pulmonary inflammation and associated loss of lung function in patients with CF (40, 41). These studies raise the prospect that the modulation of anti-inflammatory lipid mediators may be a general strategy by which P. aeruginosa regulates the host–pathogen relationship to promote the colony establishment. Despite the evidence for the involvement of these bacterial enzymes in the regulation of inflammatory lipid mediators, the CYP enzymes of P. aeruginosa have never before been examined for their ability to metabolize physiologically important fatty acids, such as arachidonic acid and its derivatives.

Here, we report the characterization of the first CYP enzyme cloned from P. aeruginosa, CYP168A1 (42, 43). Our findings indicate that CYP168A1 is a fatty acid hydroxylase with a high affinity for long-chain fatty acids, such as oleic and arachidonic acids. Additionally, CYP168A1 is capable of binding large azole inhibitors, including ketoconazole and miconazole. These results, and in silico modeling of ligands docked to the enzyme, suggest an enzyme with a large, expansive active site. Furthermore, we have characterized the hydroxylation pattern of arachidonic acid and lauric acid, a model fatty acid, establishing CYP168A1 as a medium- to long-chain fatty acid hydroxylase, attacking the carbon chain at the ω-1 and ω-2 positions. Moreover, it is capable of metabolizing arachidonic acid to 19-hydroxyeicosatetraenoic acid (19-HETE), a potent vasodilator, which may play an important role in the pathogen’s ability to colonize the mammalian lung. These results lay the groundwork for understanding the function of the CYP168A1 enzyme and its metabolic role in this important human pathogen.

Results

Expression and characterization of CYP168A1

CYP168A1 is the first CYP enzyme from P. aeruginosa PAO1 to be cloned (Fig. S1), expressed, purified, and characterized in a soluble recombinant form (42, 43). The protein was isolated and purified to homogeneity (>90%), resulting in a single band present on the denatured SDS-PAGE gel with a relative molecular weight of ~48.5 kDa (Fig. 1A). The purified protein exhibited a characteristic spectral absorption pattern for a heme containing protein, with a λmax of 417 nm for the oxidized protein and 420 nm for the sodium dithionite reduced species (Fig. 1B). Reduction of the CYP168A1 heme iron by sodium dithionite followed by carbon monoxide (CO) binding shifted the Soret peak to ~450 nm, the signature of a CYP enzyme (Fig. 1C). The Soret peak at 450 nm grew with time, ultimately reaching a plateau and stabilizing after 20 min.

Our initial expression experiment from 6 L of Escherichia coli culture yielded a total of 315 nmoles of purified and soluble recombinant CYP168A1.

Ligand binding to CYP168A1

In order to determine the chemical space occupied by ligands of CYP168A1, a series of spectral titrations were conducted with a variety of putative substrates and inhibitors, with selection criteria being based upon structural similarity to known CYP ligands. Initial experiments focused on a variety of azole compounds, which are well-known CYP inhibitors (Fig. 2). All azole ligands examined elicited a typical Type II red shift of the Soret band to ~421.5–440 nm (Fig. 2, A–D insets), reflecting direct coordination of the free electron pair of the azole ring nitrogen to the heme iron (44). The maxima and minima of these binding spectral curves were used to plot the binding isotherm (Fig. 2) and calculate the $K_d$ values for each ligand (Table 1). The four azole drugs tested...
demonstrated tight binding affinities for CYP168A1; from high to low: ketoconazole (0.68 ± 0.08 μM), miconazole (0.88 ± 0.18 μM), econazole (2.5 ± 0.4 μM), and clotrimazole (3.0 ± 0.4) (Fig. 2 and Table 1). Notably, larger azoles exhibited a higher binding affinity (lower $K_d$) than their corresponding smaller ligand counterparts, with the largest ligand—ketocono-
zole—having a submicromolar $K_d$ (Table 1). In contrast, binding of various fatty acids to the CYP168A1 ferric heme iron caused a characteristic Type I shift of the Soret band to 380 to 392 nm, reflecting displacement of the heme distal water at the sixth ligand to the heme iron and indicative of a substrate binding in the active site (Fig. 3, A–G insets). As done with the Type II azole ligands, the maxima and minima of these binding spectral curves were used to calculate the $K_d$ values for each fatty acid examined (Fig. 3 and Table 1). Consistent with our observations of larger azoles binding to the enzyme, higher-molecular-weight long-chain fatty acids were preferred over their short-chain counterparts, with the highest affinity ligands being palmitic (0.21 ± 0.04 μM), stearic (0.33 ± 0.07 μM), and oleic (0.37 ± 0.07 μM) acids (Fig. 3 and Table 1). Interestingly, while exhibiting a preference for longer-chain hydrocarbons, CYP168A1 seemed to bind both monounsaturated and polyunsaturated fatty acids with similar, submicromolar affinities (Table 1). The Type I difference spectra and tight (submicromolar, in most cases) binding affinities suggested that fatty acid ligands of chain length C10 or greater had the potential to be substrates of CYP168A1. Ligand binding was also assessed for steroids (e.g., cholesterol) and certain drugs (e.g., raloxifene, ciprofloxacin), but no spectral changes in the Soret bands were observed during the titrations.

Lauric acid metabolism by CYP168A1

The reductase electron transfer partners of CYP168A1 are currently unknown. Therefore, in order to determine if CYP168A1 was able to catalyze the oxidation of the model substrate lauric acid, initial experiments were conducted using either a series of hydroperoxide compounds as oxygen surrogates and electron donors or a spinach-derived redox partner complex consisting of the spinach ferredoxin (Fdx) and ferredoxin reductase (FdR). The spinach Fdx and FdR redox partners were previously shown to function as effective electron transfer surrogates to promote oxidation of substrates by the bacterial CYPs, CYP125A1 (45) and CYP141 (30), structurally related CYP enzymes from M. tuberculosis. Gas chromatography–mass spectrometry (GC-MS) experiments confirmed that the major metabolite produced was the 11-hydroxylauric acid (Fig. 4). The mass spectrum of the trimethylsilyl-derivatized lauric acid metabolite formed in CYP168A1 incubations with the spinach redox partners (Fig. 4B) matched the fragmentation pattern of the derivatized 11-hydroxylauric acid standard (Fig. 4A). In addition, the $m/z$ fragment of 117, corresponding to the (CH$_3$)$_3$SiO(CH$_2$-CH$_3$) ion, is a characteristic fragment of an ω-1 hydroxyl fatty acid derivative (46). Next, liquid chromatography–mass spectrometry (LC-MS) experiments were employed to quantify formation of the 11-hydroxylauric acid standard (Fig. 4A). In addition, the $m/z$ fragment of 117, corresponding to the (CH$_3$)$_3$SiO(CH$_2$-CH$_3$) ion, is a characteristic fragment of an ω-1 hydroxyl fatty acid derivative (46). Next, liquid chromatography–mass spectrometry (LC-MS) experiments were employed to quantify formation of the 11-hydroxylauric acid metabolite with the hydroperoxide compounds or the spinach redox partners. The 11- and 12-hydroxylauric acid standards were baseline separated (Fig. 5). The LC-MS trace for the lauric acid incubation of CYP168A1 with the spinach redox partners and the cofactor β-nicotinamide adenine dinucleotide phosphate (NADPH).
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Table 1
CYP168A1 binding constants for fatty acids and azoles

| Ligands       | Kd ± SE (μM) | Δ absorbance maxima (AU) |
|---------------|--------------|-------------------------|
| Fatty acids   |              |                         |
| Hexanoic acid | C6:0         | ND                      |
| Octanoic acid | C8:0         | ND                      |
| Decanoic acid | C10:0        | 21 ± 4                  |
| Lauric acid   | C12:0        | 1.9 ± 0.3               |
| Myristic acid | C14:0        | 0.83 ± 0.15             |
| Palmitic acid | C16:0        | 0.21 ± 0.04             |
| Stearic acid  | C18:0        | 0.33 ± 0.07             |
| Oleic acid    | C18:1        | 0.37 ± 0.07             |
| Arachidonic acid | C20:4      | 0.96 ± 0.07             |
| Azoles        |              |                         |
| Clotrimazole  | 3.0 ± 0.4    | 0.053                   |
| Econazole     | 2.5 ± 0.4    | 0.026                   |
| Miconazole    | 0.88 ± 0.18  | 0.036                   |
| Ketoconazole  | 0.68 ± 0.08  | 0.045                   |

Abbreviations: ND, no spectral shift detected; SE, standard error.

shows the 11-hydroxyauric acid as the major metabolite formed and presence of a minor hydroxyl metabolite matching the retention time of the 12-hydroxyauric acid, although the identity of the minor metabolite could not be confirmed by GC-MS due to its low abundance.

Hydroperoxide-driven catalysis by CYP168A1

The concentration range of the tert-butyl (tBPH) and cumene hydroperoxides (CuOOH) was tested for catalysis of lauric acid hydroxylation by CYP168A1 and formation of the 11-hydroxy derivative. As can be observed in Figure 6, maximal metabolite formation occurred following incubations of 1 μM CYP168A1 with 10 μM lauric acid for 120 min with the hydroperoxides. The results demonstrate that CYP168A1 was able to hydroxylate lauric acid utilizing the peroxide shunt pathway, bypassing the catalytic cycle necessary for transferring electrons from NAD(P)H through a CYP reductase (47). Maximal lauric acid hydroxylation activity was achieved when the concentration of each hydroperoxide was 0.25 mM (Fig. 6, A and B). However, metabolism was uniquely altered for each hydroperoxide in a concentration-dependent manner.

Redox partner-driven catalysis by CYP168A1

In order to investigate ligand metabolism requiring the transfer of electrons along the full P450 catalytic cycle, surrogates for the bacterial reductases were employed. We used the spinach Fdx and FdR as redox partners in the following experiments. The effects of varying Fdx and FdR concentrations were assessed in order to determine the optimal CYP:redox partner ratio (Fig. 7). When increasing Fdx concentrations, formation of the 11-hydroxyl metabolite was highest at the 1:20 CYP:Fdx ratio (Fig. 7A). In contrast, when increasing FdR concentrations, formation of the 11-hydroxyauric acid reached plateau at 0.2 U/ml of FdR (Fig. 7B). The optimal ratio of CYP:Fdx:FdR was determined to be 1 μM:20 μM:0.2 U/ml for maximal CYP168A1 lauric acid hydroxylation activity. However, under the 1:20 CYP:Fdx ratio conditions and the 0.2 U/ml FdR, metabolism of lauric acid exceeded the 20% substrate consumption leading to nonlinear enzyme kinetics. Thus, to allow for further assessment of CYP168A1 lauric acid hydroxylation kinetics under appropriate steady-state conditions, we set the final CYP:Fdx:FdR ratio at 1 μM:10 μM:0.05 U/ml.

Kinetics of lauric acid ω-1-hydroxylation by CYP168A1

To further elucidate the kinetic mechanism of lauric acid oxidation by CYP168A1, steady-state kinetic experiments were performed using the optimized reaction conditions described above. Due to the apparent sigmoidal nature of the data (Fig. 8), both the Michaelis–Menten and Hill fits were compared using the Akaike Information Criterion (AIC) (48). The difference of the second-order Akaike Information Criterion (AICc) is representative of the difference between the simpler model (Michaelis–Menten) minus the alternative model (Hill), which includes more fitting parameters. The difference in AICc for lauric acid ω-1-hydroxylation by CYP168A1 was determined as 16.85 (Table 2). A positive number for the difference of AICc means that the alternative model (with more parameters) has the lower AICc and is preferred. Thus, the Hill model was judged to yield a better fit for the lauric acid kinetic data set with a correct fit probability of 99.98% (Table 2). According to the Hill equation, CYP168A1 has an $S_{50}$ of 34 ± 2 μM, a $V_{max}$ of 0.12 ± 0.004 nmol/min/nmol P450, and an $n$ value, a measurement of the cooperativity of substrate binding, of 1.5 (Fig. 8 and Table 2). An $n$ value above 1 is indicative of positive cooperativity in substrate binding (49).

Arachidonic acid metabolism by CYP168A1

The low $K_d$ measured for the longer unsaturated fatty acids drove our investigation toward characterizing the potential for arachidonic acid metabolism by CYP168A1. Several precautions were taken during the arachidonic acid preparation and incubation steps to limit oxidative degradation of the unsaturated fatty acid, including working in dark conditions, preparation of new solution of arachidonic acid for each new assay, addition of antioxidants in the incubations and in the stopped solvent solution, and use of nitrogen-flushed solvents. The CYP:redox partner ratio previously optimized for lauric acid kinetic experiments was initially used to investigate arachidonic acid metabolism. LC-MS experiments were carried out with arachidonic acid and CYP168A1, and two major hydroxyl metabolites were identified, the 18-hydroxyeicosatetraenoic acid (18-HETE) and the 19-HETE (Fig. 9A). Formation of the 20-hydroxyeicosatetraenoic acid (20-HETE) metabolite was also detected, but at too low level to allow quantification (Fig. 9A). The possibility of the formation of other arachidonic acid metabolites, such as epoxyeicosatrienoic acids (EETs), also existed, but this was not examined directly as the 18-, 19-, and 20-HETE were the most abundant metabolites in our assay. Formation of the HETE metabolites was NADPH-dependent and CYP-dependent according to our control incubations. The MSMS spectra of the 18- (Figs. 9C) and 19-HETE (Fig. 9B) formed in CYP168A1 arachidonic acid incubations matched the MSMS spectra of the corresponding
Figure 3. Fatty acid binding isotherms with representative binding spectra for CYP168A1. Binding isotherms of decanoic (A), lauric (B), myristic (C), palmitic (D), stearic (E), oleic (F), and arachidonic (G) acids, with insets containing representative binding spectra obtained with 1 μM CYP168A1, were fitted to the one binding site model with R² of 0.94, 0.91, 0.88, 0.91, 0.90, 0.88, and 0.97, respectively.

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authentic standards (Fig. S2). Quantification of the 18-HETE and 19-HETE formed over time with CYP168A1 (data not shown) highlighted the requirement in optimizing the concentrations of the CYP168A1 enzyme and the spinach redox partners in the arachidonic acid incubations to achieve steady-state conditions. Effectively, the CYP:Fdx:FdR ratio had to be adjusted to 0.5 μM:5 μM:0.025 U/ml to achieve linearity in product formation up to 20 min (data not shown). Concurrently to our linearity assessment, we tested the effect of the azole ligand ketoconazole on the 18- and 19-HETE metabolite formation (Figs. 10 and S3). In the presence of 10 μM ketoconazole, CYP168A1 formation of 18- and 19-HETE was inhibited to 67.6% and 62.7% (Fig. 10), respectively, compared with the solvent control (Table 3).

Kinetics of arachidonic acid ω-1- and ω-2-hydroxylation by CYP168A1

To explore the kinetic mechanisms driving the arachidonic acid hydroxylation by CYP168A1, steady-state kinetic
experiments were performed using the optimized CYP:redox partner ratio described above for arachidonic acid. Formation of the 18- and 19-HETE metabolites was quantified using calibration standards prepared in matrix (Fig. S4). Blank matrix and carryover controls were assessed. No NADPH controls were included for each arachidonic acid concentration tested and showed no hydroxyl metabolite formation (Fig. S5). Unexpectedly, the 18- and 19-HETE formation rates measured over a range of arachidonic acid concentrations displayed substrate inhibition kinetics (Fig. 11). According to the substrate inhibition model fit of the 18- and 19-HETE formation rate data, CYP168A1 exhibits a $K_m$ of $36 \pm 18 \mu M$, $V_{max}$ of $82 \pm 33$ pmol/min/nmol P450, and $K_i$ of $16 \pm 8 \mu M$ for 18-HETE formation, and a $K_m$ of $41 \pm 25 \mu M$, $V_{max}$ of $220 \pm 110$ pmol/min/nmol P450, and $K_i$ of $13 \pm 8 \mu M$ for 19-HETE formation (Table 4). To prove that the decrease in metabolite formation is a result of substrate inhibition and not due to ion suppression coming from potential degradation contaminants from increasing concentrations of arachidonic acid, the internal standard peak areas were plotted for the standards and all incubation samples (Fig. S6). The internal standard signal was stable throughout the assay, therefore supporting that no ion suppression phenomenon impacted the HETE metabolite quantification and that the substrate inhibition observed was sustainable.

### CYP168A1 homology model

In an effort to explore the structural basis of fatty acid hydroxylation of CYP168A1, we built a homology model based on CYP107BR1, a vitamin D3 (cholecalciferol) hydroxylase from *Pseudonocardia autotrophica* (50) using USCF MODELLER v. 10.1 (51) (Fig. 12A). The model demonstrated a high-quality sequence alignment with a zDOPE score of -1.08 and a GA341 score of 1. The PyMol castp plugin calculated an active site volume of 670 Å³ with an area of 570 Å² (Fig. 12B). As observed in other CYP enzyme structures, there is the critical active site threonine (T300), which plays a role in oxygen activation, present in the long I-helix over the heme iron. Overall, the structure is relatively compact, reminiscent of other bacterial CYP enzyme structures (23, 52). Interestingly, the structure contains an abundance of aromatic residues, including 20 Phe, 8 Trp, and 8 Tyr (Fig. 12C). A number of the Phe residues appear to be participating in pi-pi stacking interactions, which may contribute to the relative stability observed with the protein, as this has also been seen with some thermostable bacterial CYP enzymes (53).

### Docking of substrates and the inhibitor ketoconazole to CYP168A1

To increase our structural understanding of how ligands interact with the enzyme, both substrates (lauric acid and...
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Figure 6. Effect of hydroperoxides on lauric acid catalysis by CYP168A1. The relative quantification of the 11-hydroxylauric acid metabolite formed by CYP168A1 with the tert-butyl hydroperoxide (A) and the cumene hydroperoxide (B) was achieved by LC-MS. Peak area ratios of the 11-hydroxylauric acid are reported as individual replicates and as means with standard deviations (error bars). Triplicate measurements were used for the mean calculations except for the 0.5 mM IBHP where only duplicate measurements were considered due to incubation error for one of the replicates. Data were analyzed by one-way ANOVA. Differences between pairwise comparisons of the specified groups are shown by *, **, *** and **** corresponding to the following significance levels: * holds p < 0.05, ** holds p < 0.01, *** holds p < 0.001, and **** holds p < 0.0001, respectively. ns, not significant.

Arachidonic acid and an inhibitor (ketoconazole) were docked to the protein using AutoDock Vina v. 1.2 (54). The most thermodynamically stable structure with the docked inhibitor, ketoconazole, with a ΔG of -8.0 kcal/mol, found it to be present in the active site in a canonical pose with the nitrogen of the azole ring 1.81 Å from the heme iron (Fig. 12D). The binding pose is similar to the CYP3A4-ketoconazole structure, with the bulk of the molecule in an extended conformation across the active site, making numerous contacts with hydrophobic residues that line the substrate access channel, including (M292, V343, V445, and V357), which likely help anchor it in a high-affinity inhibitory conformation. While the favored pose for the model substrate lauric acid found it in the active site in close proximity with many of the same hydrophobic residues (including M292, A296, V343, V445, and V357), the carbon backbone of the ligand was kinked at C7, allowing the molecule to bend and place C11 within ~5 Å of the heme iron (5.02 Å) (Fig. 12E). Additional hydrophobic contacts occur with F118, L138, and G346. However, unlike ketoconazole, we also observed contacts with important hydrophilic residues, including the catalytic threonine (T300) and S444, an active site serine residue that has a homology in CYP3A4 and is critical for oxidative activity against a number of substrates, including testosterone (55), carbamazepine (56), and diazepam (57). The most stable binding pose of lauric acid within the CYP168A1 active site returned a binding energy of ΔG of -5.7 kcal/mol. In contrast, the thermodynamically stable binding conformation of arachidonic acid found the molecule in a perpendicular orientation to the heme iron, with the carboxyl moiety extended toward the surface of the enzyme and a kink occurring at C15, positioning C18, C19, and C20 all close to the heme iron (4.72, 5.08, 5.08 Å, respectively) (Fig. 12F). The binding energy of the arachidonic acid docked structure was -7.2 kcal/mol, closer to that observed with the inhibitor ketoconazole. All of the same hydrophobic contacts observed with both ketoconazole and lauric acid were also present in the arachidonic acid structure (i.e., F118, L138, V343, G346, M292, V345). However, its larger size permitted additional contacts with residues Q80, L227, A228, M295, and A296. Again, as observed with the lauric acid structure, there was interaction between the substrate and the catalytic threonine (T300).

Discussion

CYP168A1 is the first cytochrome P450 enzyme from P. aeruginosa to be cloned, expressed, and characterized in a heterologous host (42, 43). The genome of P. aeruginosa contains four putative CYP enzymes of unknown function (10). CYP168A1 is the largest of these four putative CYP enzymes at 444 amino acids in length, significantly larger than the other three CYPs, particularly given that CYP239A1, at 386 amino acids in length, has been postulated to be a pseudogene (10).

Cytochrome P450 enzymes perform a wide variety of functions in bacteria (52), from secondary metabolite production (20–22) to carbon source utilization (15–17), to synthesis of lipids critical for cell wall integrity (52, 58). In some cases, these functions are essential to microbial growth or confer particular advantages (45); P. aeruginosa is a Gram-negative opportunistic pathogen that is particularly prevalent in the lungs of patients with cystic fibrosis, chronic obstructive pulmonary disease, and nosocomial pneumonia (59–61). In patients with compromised lung and/or immune functions, recurrent infections are common (62, 63). Successive rounds of antibiotic treatment can then lead to resistance, often mediated by the development of biofilms that make it almost impossible to eradicate the organism from the patient’s lung (6, 64). Prior studies have demonstrated that exposure to quinolones and carbapenems, antibiotics commonly used in
ICUs, is linked to the development of multidrug-resistant 
P. aeruginosa (65), reducing the therapeutic options available
to the clinician and increasing hospital mortality rates.

The UV–visible spectral data obtained from CYP168A1 are
consistent with a cytochrome P450 enzyme that is primarily
composed of holoprotein, with little P420 species present,
indicating that the protein was likely to be active (Fig. 1).

Initially, in order to determine the variety of ligand chemical
space that might be accessible to CYP168A1, we examined
several azole drugs, which are known CYP inhibitors (Fig. 2)
(66). For all the azole drugs tested, a Type II difference spec-
trum was observed, indicative of inhibitors that bind through
coordination of the azole nitrogen-free electron pair with the
heme iron (67). Interestingly, larger ligands displayed higher
affinity with the largest azole tested, ketoconazole, having the
highest affinity (0.68 ± 0.08 μM; Table 1). These results are
consistent with a rather large and expansive active site, similar
to the human CYP3A enzymes (68), that is able to accommodate
large, bulky hydrophobic ligands (Fig. 12 D). Additionally, it points to a strategy for inhibition of the enzyme
through a specific azole functionality.

The most closely related CYP enzymes to CYP168A1 that
have been thoroughly characterized are from the pathogen 
M. tuberculosis (23, 27, 66, 69). Previous studies have
demonstrated that, in the case of this obligate pathogen, 
CYP121A1 (70, 71), CYP125A1 (45, 72), and CYP142A1 (73)
are essential for bacterial survival, making them attractive drug
targets (74, 75). CYP125A1 and CYP142A1 from 
M. tuberculosis have been identi-
fied as cholesterol hydroxy-
lases, suggesting that at least some CYP enzymes from path-
ogens have evolved functions as steroid or fatty acid
hydroxylases, which are common roles for a number of mi-
crobial CYP enzymes (76, 77). This knowledge underlined our
strategy for identifying possible substrates of CYP168A1 in
order to determine its function.

Initially, we examined saturated fatty acids with chain
lengths of 6–18 carbons (Fig. 3 and Table 1). As can be
observed from the results presented in Figure 3, all fatty acid
ligands elicited a Type I difference spectrum indicative of
substrate binding, except for those of length C-8 or shorter, 
where no spectral perturbations were visible. Similar to the
trend observed with the azole compounds, CYP168A1
showed a clear preference for longer-chain fatty acids, as
determined by their $K_d$, with an optimal length of approxi-
mately 16 carbons (Table 1). This belies a large, or at least
deep, hydrophobic active site that can accommodate endog-
ogenous long-chain fatty acid substrates that may be present in
the host environment (8). Indeed, CYP168A1’s active site
volume is quite comparable with the active site volume of
CYP3A4 (520 Å$^3$) (78) and CYPBM3 (400 Å$^3$) (79) (Fig. 12 B).

Furthermore, CYP168A1 tightly bound both the mono-
unsaturated fatty acid oleic acid (0.37 ± 0.07 μM) and the
polyunsaturated fatty acid, arachidonic acid (0.96 ± 0.07 μM),
implying that both saturated and unsaturated fatty acids could
serve as substrates for CYP168A1. Interestingly, some common drugs (e.g., ciprofloxacin and raloxifene) did not exert any changes in the heme Soret spectrum, nor did the steroid cholesterol. Therefore, CYP168A1 may have a narrower substrate specificity, limited to fatty acids or similar structurally related molecules.

To determine if fatty acids could indeed be oxidized by CYP168A1, we initially examined catalysis of the model fatty acid substrate, lauric acid, using either the oxygen surrogates of tert-butyl and cumene hydroperoxides or the spinach redox partners. While CYP enzymes are readily identifiable in various bacterial genomes through their unique sequences, such as the FxxGxxxCxG heme motif (10, 80, 81), it remains a challenge to identify their active redox partners, a ferredoxin and ferredoxin reductase for a typical Type I system (67). Hence, spinach ferredoxin (Fdx) and ferredoxin reductase (FdR) have been employed as an electron delivery system in order to catalyze substrate oxidation for various bacterial CYP enzymes (30, 45). Our initial GC-MS (Fig. 4) and LC-MS (Fig. 5) experiments with the spinach redox partners confirmed the 11-hydroxylauric acid as the primary lauric acid metabolite formed, validating the identity of CYP168A1 as a subterminal fatty acid hydroxylase. In addition, according to our LC-MS analysis, a small proportion (<5%) of lauric acid was metabolized to the ω-hydroxy derivative (Fig. 5A). It is interesting to note that our docking studies determined that lauric acid was present in the active site in a somewhat unusual sterically strained conformation (Fig. 12E), which may help explain why CYP168A1 primarily performs the ω-1 oxidation over the ω. After confirming lauric acid metabolite identity, we

### Table 2

Comparison of Michaelis–Menten and Hill fits for lauric acid ω-1-hydroxylation by CYP168A1

| Model fit       | $K_m/S_{50}$ ± SE (μM) | $V_{max}$ ± SE (nmol/min/nmol P450) | $R^2$ | Correct fit probability | Difference of AICc |
|-----------------|------------------------|------------------------------------|-------|-------------------------|--------------------|
| Michaelis-Menten | 46 ± 5                 | 0.14 ± 0.006                       | 0.98  | 0.02%                  | 16.85              |
| Hill            | 34 ± 2 ($n = 1.5$)     | 0.12 ± 0.004                       | 0.99  | 99.98%                 |                    |

Abbreviations: AICc, second order Akaike Information Criterion; $n$, cooperativity value for the substrate binding to the enzyme; SE, standard error.

Figure 9. *P. aeruginosa* CYP168A1 metabolizes arachidonic acid to the 18- and 19-HETE metabolites. A, representative MRM chromatograms for arachidonic acid (10 μM) and its 18-, 19-, and 20-HETE metabolites formed in incubations with the recombinant CYP168A1 in the presence of the spinach redox partners with or without the cofactor NADPH, including the MRM chromatograms for the internal standard 20-HETE-d6. LC-MS/MS spectra for the 19-HETE (B) and 18-HETE (C) metabolites formed in arachidonic acid incubation with CYP168A1, including structure fragmentation insets.
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Figure 10. Inhibition by ketoconazole of CYP168A1 18- and 19-HETE formation. Relative quantification of the 18- and 19-HETE metabolites formed in incubations of arachidonic acid (10 μM) with CYP168A1, the spinach redox partners, and the NADPH cofactor in the presence or absence of 10 μM ketoconazole was achieved by LC-MS. Mean peak area ratios calculated from triplicate measurements are reported with the individual data points and respective standard deviations (represented graphically as error bars). Methanol was used as the solvent control. Data were analyzed by unpaired t test with exact p values indicated.

Further examined the tert-butyl (tBHP) and cumene hydroperoxide (CuOOH) concentration dependency to produce the 11-hydroxyauric acid. Both peroxide compounds equivalently stimulated catalysis of lauric acid with the optimal concentration for each peroxide used being 0.25 mM (Fig. 6), a value similar to what has been determined for other CYP enzymes exploiting the peroxide shunt pathway (47). We then sought to identify the optimal ratio of the spinach Fdx and FdR redox partners based on the hydroxylation of lauric acid to 11-hydroxyauric acid (Fig. 7). While the effect of Fdx concentration on the rate of 11-hydroxyauric acid was not saturable in our system (Fig. 7A), the concentration of FdR achieved maximal formation of metabolite at a concentration of 0.2 U/ml, indicating that FdR was the rate-limiting reactant (Fig. 7B). After adjusting the ratio of redox partners to CYP enzyme to maintain steady-state conditions, we conducted a complete kinetic characterization of lauric acid substrate oxidation by CYP168A1, including determining both the $K_m$ and $V_{max}$ values for hydroxylation to the 11-hydroxyauric acid product (Fig. 8). Somewhat surprisingly, the data best fit to the Hill equation, with an $n$ value of 1.5, $S_50$ of 34 ± 2 μM, and a $V_{max}$ of 0.12 ± 0.004 nmol/min/nmol P450, suggesting possible cooperativity in the oxidation of lauric acid. It is not unusual for CYP enzymes to exhibit cooperativity in substrate oxidation, as this has been observed with a number of bacterial (13, 82, 83) and mammalian (84–86) enzymes. However, this may be the first report of cooperative oxidation of a substrate by a CYP enzyme from a known human pathogen. While the oxidation of lauric acid defines CYP168A1 as a fatty acid hydroxylase, lauric acid is primarily a model substrate with no specific biological activity. In contrast, arachidonic acid, a tight binding ligand of CYP168A1 ($K_d = 0.96$ μM), is a well-known lipid second messenger involved in the response of cell signaling enzymes and, importantly for P. aeruginosa infection, a key regulator of inflammation (87). CYP168A1 metabolized arachidonic acid to predominantly the ω-1 19-HETE and the ω-2 18-HETE products (Fig. 9A), as confirmed by their MSMS spectra (Fig. 9, B and C) matching the authentic HETE standards (Fig. S2). It was also able to form the 20-HETE metabolite to a lesser extent (Fig. 9A). The 18-, 19-, and 20-HETE metabolites were not detected in (-) NADPH control reactions (Fig. 9A). In order to determine the potential for azole compounds to inhibit this reaction, we conducted the incubation in the presence of 10 μM ketoconazole, which significantly reduced the formation of both metabolites (Fig. 10 and Table 3), 67.6% in the case of 18-HETE and 62.7% for 19-HETE compared with the solvent control, indicating that ketoconazole is an effective in vitro inhibitor of the CYP168A1-mediated oxidation of arachidonic acid. A full kinetic analysis of arachidonic acid metabolism retrieved a $K_m$ of 36 ± 18 μM and a $V_{max}$ of 82 ± 33 pmol/min/nmol P450 for formation of the 18-HETE metabolite and a $K_m$ of 41 ± 25 μM and $V_{max}$ of 220 ± 110 pmol/min/nmol for formation of the 19-HETE (Fig. 11 and Table 4). The production of both 19-HETE and 18-HETE was subject to substrate inhibition, with negligible levels of metabolite being formed at substrate concentrations over 200 μM, and a $K_i$ of 16 ± 8 μM for 18- and 19-HETE, respectively. Thus, in comparison with the model substrate, lauric acid, which exhibited cooperativity in substrate oxidation, arachidonic acid demonstrates substrate inhibition. It should be noted that the appearance of the 18-, 19-, or 20-HETE metabolites after incubation with CYP168A1 cannot simply be due to ion suppression due to high amounts of arachidonic acid because this would also be reflected in changes in quantification of our internal standard (20-HETE-d$_6$) as well, which we do not observe (Fig. S5). While the 18-, 19-, and 20-HETE metabolites were the most abundant, it should be noted that other arachidonic acid metabolites could

Table 3

| HETE metabolites  | Methanol control (±SD) | Ketoconazole (10 μM) (±SD) | p-value | Percent inhibition |
|------------------|------------------------|---------------------------|---------|--------------------|
| 18-HETE          | 0.31 ± 0.01            | 0.10 ± 0.005              | 0.0001  | 67.6%              |
| 19-HETE          | 0.28 ± 0.02            | 0.11 ± 0.005              | 0.003   | 62.7%              |

Methanol control samples were used as the solvent control to determine the 100% CYP168A1 activity. Mean peak area ratios were calculated from triplicate measurements. Data were analyzed by unpaired t test with exact p values indicated.

Abbreviation: SD, standard deviation.
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Recently, there has been renewed focus on the physiological relevance of substrate inhibition in enzymatic systems (88, 89). It has been estimated that up to 20% of all enzyme systems may be subject to substrate inhibition (89). Far from being simply a kinetic artifact, substrate inhibition can have important physiological consequences, including but not limited to: controlling product formation, reducing buildup of toxic metabolic intermediates, or rapid termination of the signal transduction cascade (89). In terms of CYP enzymes, substrate inhibition is a well-known phenomenon that occurs often with the mammalian drug metabolizing CYPs (90). While the biological significance of the substrate inhibition observed with arachidonic acid remains unclear at this juncture, it is interesting to note that it was only detected with arachidonic acid, an inflammatory mediator occurring in the natural environment of the lung, and not with the model substrate lauric acid.

Curiously, arachidonic acid is overproduced in CF patients, including those infected with P. aeruginosa, due to a metabolic defect (36, 91). Therefore, this is a common fatty acid that the organism is likely to come into contact within the lung environment of the CF patient. Indeed, a number of studies have demonstrated the importance of arachidonic acid to the growth and/or virulence of P. aeruginosa. Rao et al. (92) established that addition of arachidonic acid and its oxidation products to P. aeruginosa cells expressing the metabolic regulator rahl led to increased biofilm formation. Additional work has shown that an increase in exposure of P. aeruginosa to arachidonic acid caused an eightfold increase in the minimum inhibitory concentration (MIC) for the antibiotic polymyxin B (93), demonstrating a link between arachidonic acid and P. aeruginosa antibiotic resistance. Furthermore, this same study revealed that P. aeruginosa will incorporate arachidonic acid into its cellular membrane when exposed to it in vitro (93), even though arachidonic acid is not synthesized natively by P. aeruginosa. In in vivo experiments using a rat model of P. aeruginosa pneumonia, it was observed that several oxidative metabolites of arachidonic acid, including 20-HETE, contributed to pulmonary vascular hyporeactivity (94). Finally, Auvin and colleagues showed that dietary supplementation of arachidonic acid in a mouse model of P. aeruginosa infection led to an increased mortality rate (95). The results from all of these studies point to an important interaction between P. aeruginosa and arachidonic acid and its metabolites.

Despite this, little is known of arachidonic acid metabolic pathways in P. aeruginosa. To date, only a single enzyme from P. aeruginosa, a secreted lipooxygenase known as LoxA, has been identified that is capable of metabolizing arachidonic acid, in this case to 15-HETE (38). Intriguingly, an early study examining metabolism of arachidonic acid derived from human blood polymorphonuclear leukocytes by P. aeruginosa produced two oxidative metabolites that were never completely characterized, but whose production was inhibited by carbon monoxide and ketoconazole (37), hallmarks of CYP metabolism and consistent with the results obtained in our study.

While it is well known that CYP metabolites of arachidonic acid, including the epoxyeicosatrienoic acids (EETs) and

also be produced, such as epoxyeicosatrienoic acids (EETs) or possibly others. In regard to their respective binding conformations in the CYP168A1 active site, arachidonic acid extends from near the heme iron to the roof of the active site cavity, significantly further than the more compact lauric acid (Fig. 12, E and F). It is likely that the larger size of arachidonic acid in comparison with lauric acid (C20 versus C12) necessitates it being extended out of the active site cavity, as it is less energetically favorable for it to adopt a more compact conformation within the active site itself. The extended conformation may allow multiple ligands to bind at high concentrations of substrate, thereby giving rise to the observed substrate inhibition.

Table 4

| HETE metabolites | $K_m \pm SE$ (μM) | $V_{max} \pm SE$ (pmol/min/nmol P450) | $K_i \pm SE$ (μM) | $R^2$ |
|------------------|------------------|----------------------------------|------------------|-------|
| 18-HETE          | 36 ± 18          | 82 ± 33                          | 16 ± 8           | 0.92  |
| 19-HETE          | 41 ± 25          | 220 ± 110                        | 13 ± 8           | 0.90  |

Abbreviation: SE, standard error.
HETEs, are not just metabolic oxidation by-products but are also important regulators of both physiological and pathophysiological processes (34, 96, 97), their production has never previously been linked to a microbial pathogen CYP enzyme. Due to the hydrophobic nature of the arachidonic acid metabolites, they often tend to accumulate with tissue lipids (98), but upon stimulation with hormones, the metabolites are released to act either through paracrine or autocrine pathways. In vitro experiments have demonstrated that multiple mammalian CYP enzymes, including CYP2E1, CYP2J9, and CYP2U1, can metabolize arachidonic acid to both the 19- and 20-HETE oxidation products (99–101). In mammals, 19-HETE acts as a potent vasodilator, renal Na⁺-K⁺ ATPase activator, and platelet aggregation inhibitor (102–104). While it is currently unclear what advantage vasodilation would confer to *P. aeruginosa*, it is relevant to the modulation of immune response as it is an important aspect of inflammation. Vasodilation increases blood flow to the affected area and also increases the permeability of the blood vessel walls. Whereas 20-HETE, the minor CYP168A1 metabolic product, is known to be important in the regulation of vascular tone and blood flow, as well as playing a role in inflammation by stimulating the production of various proinflammatory mediators, including: PGE2, cytokine tissue necrosis factor alpha (TNFα), and the chemokines IL-8, IL-12, IL-14 (105). The physiological role of 18-HETE is less well defined, but it may also be

![Figure 12. Docking of ligands to the CYP168A1 homology model. A, stylized amino acid backbone of the CYP168A1 homology model, with the heme prosthetic group shown in red (in all structures), (B) the P-cast defined active site of the enzyme (represented by magenta balls), (C) aromatic residues of CYP168A1 (shown in cyan), (D) active site cutaway of the docked structure of ketoconazole (cyan) within the CYP168A1 active site, showing the azole nitrogen in close proximity to the heme iron, (E) active site cutaway of docked structure of lauric acid (magenta); for simplicity sake, only the carbon skeleton backbone is shown; note the kinked and extended structure of the substrate, (F) active site cutaway of the docked structure of arachidonic acid (magenta) within the CYP168A1 active site; the conformation of the substrate is similarly kinked and extended as in the case of lauric acid. All figures were generated using UCSF Chimera (https://www.cgl.ucsf.edu/chimera/).](image-url)
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important in the regulation of blood flow and blood vessel contractility (106).

*P. aeruginosa* is an opportunistic pathogen and can exist as an innocuous soil microorganism in the natural environment (107). Indeed, recent work examining the potential of *P. aeruginosa* as a bioremediation vector has demonstrated that its complement of CYP enzymes is capable of oxidizing medium to short-chain alkanes (32). It is possible that the *P. aeruginosa* CYPs may have originally evolved to utilize carbon sources readily available in the local environment, such as lauric acid.

Only more recently, evolutionary speaking, has *P. aeruginosa* evolved the ability to adapt to the mammalian lung (108). As a consequence, its complement of enzymes may now be adjusting to new roles. In general terms, the adaption of *P. aeruginosa* CYP168A1 to metabolize arachidonic acid to 18-, 19-, and nominally 20-HETE may reflect a burgeoning ability of the pathogen to modulate the immune response of the host organism in order to make the lung environment more palatable for colonization. A hallmark of pathogen success as a parasite is the ability for it to be able to “communicate” with the host organism through the production of proteins and small-molecule metabolites that can modulate the immune response and/or improve the characteristics of the host environment in order to permit pathogen growth and replication (109, 110). *P. aeruginosa* may accomplish this through its metabolism of arachidonic acid to metabolites, such as 18-, 19-, and 20-HETE, that provide important physiological functions. There are other possibilities to be considered as to why arachidonic acid supplementation might alter host immune response, including differential oxylipin profiles or effects on bacterial growth rate that are not directly related to CYP168A1 metabolism. Furthermore, an additional intriguing possibility may be that 19-HETE is an anti-inflammatory proresolving mediator that downregulates macrophage function in the host (111). Finally, it is also possible that CYP168A1 may hydroxylate omega-3 fatty acids to produce more potent oxylipin mediators. While the exact role of the production of these metabolites by *P. aeruginosa* remains a mystery, it is an area of active investigation in our laboratory, and it may point toward a critical role for *P. aeruginosa* CYP168A1 in the maintenance of infection.

In summary, we have characterized the first CYP enzyme from *P. aeruginosa* as a fatty acid hydroxylase capable of metabolizing arachidonic acid to 18-, 19-, and nominally 20-HETE, all important physiological mediators. Further investigation of the role that this enzyme plays in the pathogen life cycle may possibly reveal new insights on its ability to grow and replicate in the mammalian lung and also new metabolic functions.

Experimental procedures

**Materials**

Lauric, decanoic and stearic acids, and 5-aminolevulinic acid hydrochloride were purchased from Acros Organics (Fair Lawn, NJ). The 11-hydroxylauric and 12-hydroxylauric-d20 acid standards were purchased from Santa Cruz Biotechnology. The 12-hydroxyauric, palmitic and arachidonic (oil) acids, clotrimazole, imidazole, tert-butyl (tBPH), and cumene hydroperoxides (CuOOH) were obtained from Sigma-Aldrich (St Louis, MO). Myristic acid and econazole were purchased at VWR International, and ketoconazole was from Toronto Research Chemicals. Oleic acid and miconazole were obtained from Thermo Fisher Scientific. Ampicillin, arachidonic acid sodium salt, 18-HETE, 19(S)-HETE, 20-HETE, and 20-HETE-d6 were all purchased from Cayman Chemical. Isopropyl-β-D-1-thiogalactopyranoside (IPTG), phenylmethanesulfonl fluoride (PMSF), glucose-6-phosphate, and β-nicotinamide adenine dinucleotide phosphate (NADP+) were obtained from Alfa Aesar. Glucose-6-phosphate dehydrogenase, the spinach Fdx and FDR were purchased from Sigma-Aldrich. All other chemicals and solvents were obtained from standard suppliers and were of reagent or analytical grade.

**Construction of CYP168A1 expression vector and expression of the recombinant CYP168A1 protein**

The National Center for Biotechnology Information (NCBI) amino acid sequence NP_251165 is a 444 amino acid sequence classified as a putative CYP from *P. aeruginosa* PAO1 (112) and designated according to the P450 nomenclature as CYP168A1 (10). The amino acid sequence was initially reverse translated to DNA using the Sequence Manipulation Suite website (113). CYP168A1 cDNA sequence was then codon optimized for expression in *E. coli* using the GenScript GenSmart Codon Optimization Tool (https://www.genscript.com/gensmart-free-gene-codon-optimization.html) (Fig. S1A). Finally, the codon-optimized DNA sequence was engineered with four histidine residues at the 3’-end of the sequence prior to the stop codon and inserted into a pUC57 vector using Ndel and HindIII engineered restriction site sequences at the 5’ and 3’ ends of the CYP168A1 DNA coding sequence, respectively. Following transformation of *E. coli* DH5α cells (Invitrogen), pUC57-CYP168A1 plasmids were isolated using the Qiagen Miniprep Kit (Qiagen). The CYP168A1-optimized cDNA insert was removed from the plasmid using Ndel and HindIII restriction enzymes and isolated using agarose gel electrophoresis and the Qiagen Gel Extraction Kit (Qiagen) for ligation into a similarly digested and isolated pCWori+ CYP expression vector (114). This plasmid, designated as pCWori-CYP168A1, was then used to transform *E. coli* DH5α cells in preparation for expression of the CYP168A1 protein (Fig. S1B).

CYP168A1 was expressed under the control of the tac promoter of the pCWori+ plasmid using *E. coli*-DH5α F′IQ cells (Thermo Fisher Scientific) in Terrific Broth (TB) medium. Briefly, 10 ml of an overnight pCWori-CYP168A1 *E. coli*-DH5α starter culture consisting of Luria-Bertani (LB) medium supplemented with 200 μg/ml ampicillin was used to inoculate each liter of TB (also containing 200 μg/ml ampicillin). The bacterial culture was incubated at 37 °C under agitation (250 rpm) until the optical density reached 0.5 to 0.8 at 600 nm. Then IPTG and 5-aminolevulinic acid were added at
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0.5 and 0.25 mM final concentration, respectively. The expression culture was allowed to grow for another 24 h at 25 °C and 180 rpm agitation. The bacterial cells were then pelleted by centrifugation at 3400 g for 45 min, and the cell pellets were stored at −80 °C until purification of the expressed CYP168A1 enzyme.

Purification of the (His)-Tagged CYP168A1

Expressed CYP168A1 was purified using fast protein liquid chromatography with a HiTrap-HP affinity column (GE Healthcare). The bacterial cell pellets were thawed on ice and resuspended in buffer A consisting of 50 mM Tris-HCl pH 7.5, 50 mM NaCl, 0.1 mM ethylenediaminetetraacetic acid (EDTA), 20 mM imidazole, and 1 mM PMSF. About 4 ml of buffer A was used for resuspension of a gram of bacterial cell pellet. After addition of lysozyme (0.3 mg/ml) and DNase (700 U), the bacterial cell suspension was stirred on ice for 30 min. Cells were then lysed on ice using a Branson sonicator set at 50% power and three 4-min bursts with 2 min resting time between each burst. Following cell lysis, whole cells and cell debris were separated by ultracentrifugation at 100,000 g for 60 min.

The ultracentrifugation supernatant containing the recombinant CYP168A1 protein was loaded onto a HiTrap-HP column (5 ml) previously equilibrated with buffer A. The column was subsequently washed with five column volumes of buffer A. CYP168A1 was then eluted with a gradient of imidazole using the elution buffer B (50 mM Tris-HCl, 0.1 mM EDTA, 0.2 M imidazole). Red-colored fractions (5 μl) were analyzed for purity by SDS-PAGE gel electrophoresis on a 4 to 15% Mini-Protein TGX precast gel in a Tris/Glycine/SDS buffer. The proteins on the SDS-PAGE gel were revealed with the Bio-Rad Bio-Safe Coomassie G-250 stain. Fractions containing the bulk of the recombinant CYP168A1 protein were pooled and then dialyzed at 4 °C in 50 mM Tris-HCl pH 7.5, 0.1 mM EDTA, and 0.1 mM dithiothreitol. The protein concentration of the purified CYP168A1 was determined by bicinchoninic acid (BCA) assay (Pierce, Thermo Fisher Scientific), and the final concentration of the ferrous-CO protein was determined using UV–visible spectroscopy, with an extinction coefficient of ε = 91 mM⁻¹ cm⁻¹ at the wavelength of 450 nm (115). UV–visible spectroscopy was further used to characterize the spectral absorption pattern of the absolute and reduced CYP168A1 protein.

Ligand Kd determination by optical difference spectroscopy

To determine ligand selectivity of CYP168A1, UV–visible difference spectra were acquired on a Varian Cary 50 Bio UV–visible scanning spectrophotometer (Agilent) for various ligands, including antifungal azole compounds and fatty acids. Both sample and reference chambers contained 1 ml of 1 μM CYP168A1 in 100 mM potassium phosphate, pH 7.4. Prior to initiating the titration, a baseline was recorded between 350 and 500 nm. Aliquots of ligand stock solutions prepared by serial dilution in dimethyl sulfoxide (DMSO) were added to the sample cuvette, whereas equal volume of vehicle solvent was added to the reference cuvette to determine the difference spectrum at varying concentrations. The absolute changes in absorbance deriving from a minimum of triplicate titrations were plotted as a function of ligand concentration and fitted to the one binding site model using the GraphPad Prism software (version 9.0.0, GraphPad software).

Recombinant CYP168A1 lauric acid in vitro metabolic assays

Due to its high affinity for CYP168A1 and known activity as a substrate for other microbial CYP enzymes (53, 77), lauric acid was used as a model substrate for metabolism studies. For the hydroperoxide-driven catalysis, solutions of tbPh and CuOOH were freshly prepared in DMSO and incubated at different concentrations up to 120 min with 1 μM CYP168A1 and 10 μM lauric acid in 100 mM potassium phosphate, pH 7.4 (47). Reactions were stopped by the addition of an equal volume of ice-cold methanol containing 60 ng/ml 12-hydroxylauric-d20 acid as internal standard. Samples were centrifuged at 2500 g and 4 °C for 20 min for protein precipitation. Supernatants were transferred to high-performance liquid chromatography (HPLC) vials, and aliquots of 5 μl were analyzed by LC-MS. For the redox partner-driven catalysis, different concentrations of spinach Fdx and FdR were assessed to obtain the optimal CYP:redox partner ratio. Initial linearity experiments were done establishing linearity up to 30 min. The incubations with the spinach redox partners (200 μl) were carried out in 100 mM potassium phosphate, pH 7.4, and 3 mM MgCl2 with 1 μM CYP168A1 and 10 μM lauric acid. Concentrations of spinach Fdx and FdR were between 2 and 20 μM and 0.05 to 0.3 U/ml, respectively. After an equilibration at 37 °C for 3 min, the reactions, prepared in triplicate, were initiated by the addition of a NADPH-regenerating system mix consisting of NADP⁺ (1 mM), D-glucose-6-phosphate (10 mM) and glucose-6-phosphate dehydrogenase (2 IU/ml). The reactions were incubated for 30 min at 37 °C under agitation and were stopped by the addition of ice-cold methanol (200 μl) containing 60 ng/ml 12-hydroxylauric-d20 acid internal standard. Incubations without the NADPH-regenerating system mix served as negative controls. Precipitated proteins were collected by centrifugation of the stopped reaction samples for 20 min at 2500 g and 4 °C. Supernatants were transferred to HPLC vials, and aliquots of 5 μl were analyzed by LC-MS. The 11-hydroxylauric acid metabolite was quantified based on a calibration curve prepared in matrix and ranging from 0.1 μM to 10 μM. Blank matrix and standard zero samples were included in the LC-MS analysis. The matrix blank was made up of buffer solution (consisting of 100 mM potassium phosphate, pH 7.4, 3 mM MgCl2), CYP168A1, Fdx, FdR, and methanol. The standard zero was prepared similarly than the blank matrix, except the methanol also contained 60 ng/ml 12-hydroxylauric-d20 acid internal standard.

For the kinetic reactions, similar assay conditions were used with concentrations of lauric acid ranging from 2.5 to 250 μM. To ensure steady-state kinetic conditions and less than 20% substrate depletion, the concentration of CYP enzyme, Fdx,
and FdR used was 1 μM, 10 μM, and 0.05 U/ml, respectively. The 11-hydroxylauric acid metabolite was quantified based on a calibration curve prepared in matrix and ranging from 0.1 μM to 10 μM and included matrix blank and standard zero samples as described above. The mean metabolite formation rate values obtained from triplicate determinations were fit to the Michaelis–Menten (hyperbolic) and Hill (sigmoidal) equations using GraphPad Prism software (version 9.0.0). Comparison of the best fit was based on the second-order AICc analysis (Table 2).

**Lauric acid metabolite analysis by GC-MS**

Lauric acid metabolites generated in incubations of 100 μM lauric acid with CYP168A1 (5 μM) and the spinach redox partners Fdx (5 μM) and FdR (0.13 U/ml) were extracted twice with dichloromethane and dried at room temperature under nitrogen flow. Samples, resuspended in acetonitrile, were derivatized by addition of 50 μl N-methyl-N-trimethylsilyl-trifluoroacetamide (containing trimethylchlorosilane at 1% v/v) followed by 20 min incubation at 70 °C. Derivatized samples were transferred to Teflon capped vials for GC-MS analysis on an Agilent Technologies 5977/78900 gas chromatograph using an Agilent HP-5MS column (30 m × 0.25 mm inside diameter × 0.25 μm). Separation of trimethylsilyl-derivatized lauric acid and hydroxyl metabolites was achieved by temperature gradient as follows: 70 °C for 1 min, increased 25 °C/min up to 170 °C, then increased 5 °C/min up to 200 °C and increased 20 °C/min up to 280 °C, held for 5 min at 280 °C. GC parameters were as follows: inlet temperature of 250 °C, splitless constant flow mode at 7.57 ml/min, and MS transfer line at 230 °C. Lauric acid and its metabolites were ionized using electron impact ionization and detected by a single quadrupole in scan mode from 30 to 500 mass units. MS parameters were as follows: source temperature of 230 °C, quadrupole temperature of 150 °C. The GC-MS system was controlled by an Agilent MassHunter Workstation. Data were analyzed using Agilent MassHunter Quantitative Analysis. Identification of the trimethylsilyl-derivatized 11-hydroxylic acid was confirmed using an authentic standard and verified using the NIST17 GC-MS mass spectral database (116).

**LC-MS method for lauric acid hydroxylation**

The lauric acid incubation samples with the recombinant CYP168A1 enzyme were analyzed by LC-MS with a Waters Acquity Ultra-Performance Liquid Chromatography (UPLC) system interfaced by electrospray ionization with a Waters Xevo TQ-S micro tandem quadrupole mass spectrometer (Waters Corp.) in negative ionization mode and with multiple reaction monitoring (MRM) scan type. Due to limited fragmentation of lauric acid, its metabolites, and the internal standard, a parent-to-parent mass transition strategy was employed. The following mass transitions, collision energies (CEs), and cone voltages (CVs) were used to detect the respective analytes: 199.1 > 199.1, CE = 10 V, CV = 40 V for lauric acid, 215.1 > 215.1, CE = 10 V, CV = 40 V for the hydroxylauric acid metabolites and 235.2 > 235.2, CE = 8 V, CV = 20 V for the internal standard 12-hydroxylauric-d20 acid. The following source conditions were applied: 1 kV for the capillary voltage, 150 °C for the source temperature, 500 °C for the desolvation temperature, and 900 l/h for the desolvation gas flow. Lauric acid and its hydroxylated metabolites were separated on a Waters BEH C18 column (1.7 μm, 2.1 × 50 mm) by flowing 2 mM ammonium acetate in water and in methanol at 0.4 ml/min. The following gradient was used: 10% organic (methanol) held for 0.5 min, increased first to 45% over 0.5 min, then increased to 53% over 2 min, and finally increased to 98% over 0.2 min and held at 98% over 1.8 min. To limit soiling of the source, a divert directing the LC flow to waste was set at 3.5 min before elution of lauric acid. The MS peaks were integrated using QuanLynx software (version 4.1, Waters Corp.), and the analyte/internal standard peak area ratios were used for relative quantification. For determination of the hydroxy metabolite concentration, the regression fit was based on the analyte/internal standard peak area ratios calculated from the calibration standards, and the analyte concentration in the incubations was back-calculated using the weighted (1/x) linear least-squares regression.

**Recombinant CYP168A1 arachidonic acid in vitro metabolic assays**

The optimized CYP:Fdx:FdR ratio used for lauric acid kinetic experiments was initially used to investigate arachidonic acid metabolism and assess linearity of the reaction over 60 min. However, under these conditions, metabolite linearity could not be established, and decrease in the CYP168A1, Fdx, and FdR concentrations was necessary. The CYP:Fdx:FdR ratio of 0.5 μM:5 μM:0.025 U/ml allowed achievement of product formation linearity up to 20 min in order to remain under steady-state conditions. Incubations of arachidonic acid with CYP168A1 and the spinach redox partners were done in dim light conditions and with slight modifications of McDonald et al. (117). Briefly, arachidonic acid (sodium salt; Cayman Chemical), freshly prepared in nitrogen-flushed methanol to yield final concentrations between 2.5 and 200 μM in the reactions, was incubated with CYP168A1 and the spinach redox partners in 100 mM potassium phosphate (pH 7.4), 3 mM MgCl2, and 1 mM sodium pyruvate. After an equilibration at 37 °C for 3 min, the reactions, prepared in triplicate, were initiated by the addition of a NADPH-regenerating system mix consisting of NADP+ (1 mM), D-glucose-6-phosphate (10 mM) and glucose-6-phosphate dehydrogenase (2 IU/ml). After 20 min at 37 °C and under agitation, the enzymatic reactions were stopped by the addition of ice-cold nitrogen-flushed 4:1 acetonitrile:methanol (200 μl) containing 600 ng/ml 20-HETE-d6 internal standard and 0.02% 2,6-di-tert-butyl-4-methylphenol. Incubations without the NADPH-regenerating system mix served as negative controls. Furthermore, additional incubations were carried out without any CYP168A1 enzyme present to ascertain that formation of the hydroxyl metabolites was dependent on the presence of the cytochrome P450 enzyme. For the arachidonic acid incubations with ketoconazole, a
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Ketoconazole stock solution was made in methanol and was added in the CYP168A1 reactions prepared in triplicate to yield a final concentration of 10 μM ketoconazole. Methanol solvent control incubations were done in parallel. Precipitated proteins were collected by centrifugation of the stopped reaction samples at 2500g and 4 °C for 20 min. Supernatants were transferred to HPLC vials, and aliquots of 5 μl were analyzed by LC-MS. The 18-HETE and 19-HETE metabolites were quantified based on calibration curves prepared in matrix and ranging from 0.05 μM to 10 μM and from 0.1 μM to 10 μM, respectively (Fig. S4). Blank matrix and standard 0 samples were included in the LC-MS analysis (Fig. S4). The matrix blank was made up of buffer solution (consisting of 100 mM potassium phosphate, pH 7.4, 3 mM MgCl₂, 1 mM sodium pyruvate, CYP168A1, Fdx, FdR, and 4:1 acetone-trile: methanol (containing 0.02% 2,6-di-tert-butyl-4-methylphenol). The standard zero was prepared similarly to the blank matrix, except for the 4:1 acetone-trile: methanol containing 600 ng/ml 20-HETE-d₆ internal standard. The 19(S)-HETE standard was used to prepare the calibration curve, since no racemic mixture of 19-HETE was commercially available. For the kinetic experiment, the mean metabolite formation rate values obtained from triplicate determinations were fit to the substrate inhibition model using GraphPad Prism software (version 9.0.0).

LC-MS method for arachidonic acid hydroxylation

The arachidonic acid incubation samples with the recombinant CYP168A1 enzyme were analyzed by LC-MS with a Waters Acquity Ultra-Performance Liquid Chromatography (UPLC) system interfaced by electrospray ionization with a Waters Xevo TQ-S micro tandem quadrupole mass spectrometer (Waters Corp.) in negative ionization mode and with multiple reaction monitoring (MRM) scan type. The following source conditions were applied: 1 kV for the capillary voltage, 150 °C for the source temperature, 500 °C for the desolvation temperature, and 900 l/h for the desolvation gas flow. The following mass transitions, CEs, and CVs were used to detect the respective analytes: 303.1 > 259.1, CE = 12 V, CV = 20 V for arachidonic acid, 319.0 > 261.3, CE = 18 V, CV = 20 V for the 18-HETE, 319.0 > 231.3, CE = 15 V, CV = 20 V for the 19-HETE, 319.0 > 289.3, CE = 15 V, CV = 20 V for the 20-HETE, and 325.1 > 281.2, CE = 15 V, CV = 20 V for the internal standard 20-HETE-d₆. Arachidonic acid and its hydroxylated metabolites were separated on a Waters BEH C18 column (1.7 μm, 2.1 x 100 mm) by flowing 2 mM ammonium acetate in water and in 4:1 acetone-trile: methanol at 0.3 ml/min. The following gradient was used: 55% organic (4:1 acetone-trile: methanol) held for 3.5 min, increased to 98% over 0.5 min, and held at 98% over 2 min. The MS peaks were integrated using QuanLynx software (version 4.1, Waters Corp.), and the analyte/internal standard peak area ratios were used for relative quantification. For determination of the hydroxy metabolite concentration, the regression fit was based on the analyte/internal standard peak area ratios calculated from the calibration standards, and the analyte concentration in the incubations was back-calculated using the weighted (1/x) linear least-squares regression. For acquisition of the HETE metabolite MSMS spectra, daughter scans of m/z 319 were acquired in centroid mode between m/z of 50 and 350 using the above source conditions and a CE of 18 V.

CYP168A1 homology model construction

A BlastP search of the CYP168A1 sequence (NP_251165) against sequences in the Protein Data Bank (https://www.rcsb.org/) revealed that the top scoring hit is a protein known as CalO2, a putative CYP enzyme from Micromonospora echinospora, generating a MAX score of 145. However, this organism is not closely related to P. aeruginosa phylogenetically. Therefore, the more closely related fourth top scoring hit CYP Vdh (CYP107BR1), a vitamin D hydroxylase from Pseudomonas autotrophic, was used as a template for building the homology model of CYP168A1. Our choice for model template was based both on sequence identity and enzyme function, since all known CYPs have the same general protein fold. In this case, CYP107BR1 was chosen, in part, because it is a vitamin D hydroxylase, with similarity to fatty acid hydroxylases. Both CYP168A1 and CYP107BR1 were used as target and template, respectively. Due to the lack of similarity of the first 21 amino acids to any known sequence in the PDB or NCBI RefSeq database, these residues were omitted for the purposes of homology model construction. Removal of these residues resulted in greater than >90% sequence coverage and 34% sequence identity between CYP168A1 and CYP107BR1.

Models were generated using UCSF MODELLER v. 10.1 (51) ran locally via the UCSF Chimera v. 1.15 GUI interface (118). MODELLER parameters included an output of five independent models, nonwater HETATM residues from the template (heme), and hydrogen atoms. The final model chosen produced a zDOPE score, an atomic distance-dependent statistical score where negative values indicate better models of -0.2. Additionally, the model had a GA341 score of 1. The GA341 score is a model score derived from statistical potentials where a value >0.7 generally indicates a reliable model, i.e., >95% probability of having the correct fold. Following initial model generation, unstructured loops were refined using the loops refinement MODELLER v. 10.1 plugin in Chimera. After loop refinement, the zDOPE score was reduced to -1.08, and the GA341 score remained unchanged. This final refined model was used for all subsequent ligand docking studies.

Docking of substrates and ketoconazole to CYP168A1

In order to understand how substrate and inhibitor ligands structurally interact with CYP168A1, an in silico docking study was undertaken using AutoDock Vina v. 1.2 (54) and the CYP168A1 homology model as the receptor template. The protein was prepared for docking using MGLTools, AutoDock Tools v. 1.5.7 (The Scripps Research Institute) by adding polar hydrogens and assigning partial charges. Coordinates for the docking grid search space were established by defining the enzyme active site, with the final parameters being: grid box center; x-center = 9.285, y-center = 21.861, z-center = 19.061,
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and the total number of grid points in each dimension being: x-dimension = 24.664, y-dimension = 29.966, and z-dimension = 29.793. The ligands selected for docking were ketoc- nazole, a Type II inhibitor, and the fatty acid substrates, lauric acid and arachidonic acid. Each ligand file was downloaded from the Protein Data Bank (PDB: https://www.rcsb.org/) and parameterized for docking in the following manner: 1) addition of polar hydrogens, 2) assessment (and assignment, when necessary) of rotatable bonds, and 3) assignment of partial charges. Both receptor (protein) and ligand files were saved in the PDBQT format. A configuration file docking script was prepared in simple text format with the energy range set to 4 and the exhaustiveness search parameter set to 8. AutoDock Vina v. 1.2 was invoked using the configuration file and PDBQT.out and log-out files. Output files were qualitatively and quantitatively analyzed by using the ViewDock function of UCSF Chimera and ranked by binding energy (ΔG). The most energetically favorable binding mode for each ligand was reported as 0 rmsd. All figures were generated using UCSF Chimera v1.15 (118).

Data availability

All raw and processed data may be obtained by contacting the corresponding author directly.

Supporting information—This article contains supporting information. Figs. S1–S7 are available in the Supporting Information document and include the codon-optimized cDNA sequence, LC-MS data for the arachidonic acid assays with no NADPH and blank matrix controls, calibration standards, and internal standard peak areas for the kinetic samples and the BLASTp protein sequence alignment of CYP168A1 with CYP107BR1. The homology model for CYP168A1 used in this study is made available as Supporting Information (Supplemental Data).

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Abbreviations—The abbreviations used are: 15-epi LX44, 15-epi lipoxin A4; AICc, Akaike information criterion; BCA, bicinchoninic acid; CE, collision energy; CF, cystic fibrosis; CO, carbon monoxide; CuOOH, cumene hydroperoxide; CV, cone voltage; CYP, cytochrome P450; DMSO, dimethyl sulfoxide; EDTA, ethylenediaminetetraacetic acid; EET, epoxyeicosatrienoic acids; FdR, ferredoxin reductase; Fdx, ferredoxin; GC-MS, gas chromatography–mass spectrometry; HETE, hydroxyeicosatetraenoic acid; HPLC, high-performance liquid chromatography; IPTG, isopropyl-β-D-1-thiogalactopyranoside; IS, internal standard; LB, Luria–Bertani; LC-MS, liquid chromatography–mass spectrometry; MIC, minimum inhibitory concentration; MM, multiple reaction monitoring; MSMS, tandem MS; NADPH, β-nicotinamide adenine dinucleotide phosphate (reduced); NCBI, National Center for Biotechnology Information; PMSF, phenylmethylsulfonylfluoride; TB, tert-butyl hydroperoxide; UPLC, ultraperformance liquid chromatography.

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