Modes of heme binding and substrate access for cytochrome P450 CYP74A revealed by crystal structures of allene oxide synthase

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Edited by Richard A. Dixon, The Samuel Roberts Noble Foundation, Ardmore, OK, and approved June 23, 2008 (received for review April 28, 2008)

Cytochrome P450s exist ubiquitously in all organisms and are involved in many biological processes. Allen oxide synthase (AOS) is a P450 enzyme that plays a key role in the biosynthesis of oxylipin jasmonates, which are involved in signal and defense reactions in higher plants. The crystal structures of guayule (Parthenium argentatum) AOS (CYP74A2) and its complex with the substrate analog 13(S)-hydroxyoctadeca-9,11E-dienoic acid have been determined. The structures exhibit a classic P450 fold but possess a heme-binding mode with an unusually long heme-binding loop and a unique I-helix. The structures also reveal two channels through which substrate and product may access and leave the active site. The entrances are defined by a loop between β3-2 and β3-3. Asn-276 in the substrate binding site may interact with the substrate's hydroperoxy group and play an important role in catalysis, and Lys-282 at the entrance may control substrate access and binding. These studies provide both structural insights into AOS and related P450s and a structural basis to understand the distinct reaction mechanism.

oxylipin | jasmonate | guayule

Cytochrome P450s are one of the largest superfamilies of enzymes and are ubiquitously distributed in all biological organisms. They are heme-containing enzymes catalyzing a wide range of chemical reactions (1, 2). P450s play very important roles in drug metabolism and detoxification for humans and animals and are key players in the synthesis of natural products, such as antibiotics in microorganisms and a broad range of secondary metabolites in plants. Allen oxide synthase (AOS) is a P450 enzyme that plays a key role in the biosynthesis of oxylipins, bioactive compounds involved in signal and defense reactions in higher plants, mammals, and algae (3, 4). Jasmonates are plant oxylipins and ubiquitous plant growth regulators and are synthesized via the AOS branch of the lipoxygenase (LOX) pathway. AOS catalyzes a dehydration reaction to convert 13(S)-hydroperoxide (Fig. 1), derived from linolenic acid by LOX, to allene oxide, which is further cyclized by allene oxide cyclase, leading to the formation of jasmonic acid. Structures of Arabidopsis thaliana allene oxide cyclase and some LOXs have been reported (5, 6), but no AOS structure has been characterized to date.

Guayule (Parthenium argentatum) can accumulate an extraordinarily large number of rubber particles. AOS from guayule is the most abundant protein associated with rubber particles. It compromises ∼50% of the proteins extracted from guayule rubber particles, which was found to be a P450 enzyme with AOS activity (7), similar to the AOS identified from flaxseed (8, 9). However, the possible involvement of such an active enzyme in rubber biosynthesis remains unclear.

Although most AOSs from plants are membrane-associated, AOSs from guayule and corn, and recombinant AOSs from barley and tomato, have no N-terminal membrane anchor and are cytosolic (9, 10). A distinctive feature of guayule AOS is the absence of the putative transit sequence, which is present in flaxseed AOS and others for targeting to plastids or mitochondria.

AOSs are peroxide-metabolizing P450s and belong to the CYP74A group. They are atypical P450s, are self-sufficient, and do not require molecular oxygen and NADPH reductase. AOSs use an acyl hydroperoxide both as the substrate and the oxygen donor. The typical class I and II P450s require electron transfer partners, i.e., FAD-containing reductase and an iron sulfur reductase (for class I), and a P450 reductase (for class II). There is another class of P450s that receive electrons directly from NAD(P)H (2). The biochemical mechanisms underlying P450 reactions, especially in biosynthesis, are not well understood.

AOS from guayule has been classified as CYP74A2 with enzymatic activity toward 13(S)-hydroperoxide (7) but not 9(S)-hydroperoxide (Z.P., unpublished results). All CYP74A subfamily enzymes use 13(S)-hydroperoxides as substrates and are named 13-AOS (11). There are two other types of AOSs with different substrate specificities. The 9/13-AOS enzymes from barley and rice recognize both 9(S)-hydroperoxides and 13(S)-hydroperoxides (12). The 9-AOSs from tomato and potato have specificity for 9(S)-hydroperoxides (13, 14). The 9/13- and 9-AOSs are grouped in the CYP74C subfamily.

Structural studies of P450s provide an essential basis for understanding their complex catalytic reactions. To date, many structures have been reported for soluble bacterial/mitochondrial class I P450s (15–21). Class II enzymes in eukaryotes are microsomal P450s containing a membrane-binding anchor at the N terminus. Several mammalian microsomal P450 structures have been solved, including the closely related rabbit P450s 2C5 and 2B4, and the human P450s 2C8, 2C9, 2A6, 3A4, and 2D6 (22–29). Structures have also been reported for other classes of P450s, e.g., the fungal nitric oxide reductase P450nor (30), and prostacyclin synthases (PGIS, CYP8A1), which are involved in endoperoxide metabolism. To catalyze the isomerization of prostaglandin H2 to prostacyclin (31, 32). A very large number of P450s are present in plants, with nearly 2,900 named plant P450s in the...
P450 database (http://drnelson.utmem.edu/CytochromeP450.html). Many are known to be involved in the biosynthesis of natural products. So far, no plant P450 structure has been reported.

Based on amino acid sequence information, AOS contains an unusual heme-binding region and possesses a defective I-helix without the highly conserved threonine, suggesting an altered I-helix function (7). Homology modeling of AOSs is difficult because of the very poor sequence identity with any other P450 enzymes. Detailed structural information is therefore essential to understand the catalytic mechanism of AOS. Recently we successfully expressed, purified, and crystallized AOS (33).

Here, we report crystal structures of AOS (CYP74A2) from guayule (P. argentatum) and its complex with the substrate analog 13(S)-hydroxyoctadeca-9z,11E-dienoic acid [13(S)-HODE]. These plant P450 crystal structures revealed distinct structural features of AOS and its interactions with heme and substrate analog. From the structures, a heme-binding mode and two access channels for substrate to enter and/or for product to leave the active site have been identified, along with residues important for substrate binding and catalysis.

Results and Discussion

Structure Determination and Overall Structure. The crystal structures of AOS in native form (Native 1) and in complex with the substrate analog 13(S)-HODE were determined by using the multiple isomorphous replacement with anomalous scattering (MIRAS) method. Another AOS native crystal form with a small cell unit, denoted as Native 2, diffracted to 1.8Å resolution, and the structure was determined by using the molecular replacement (MR) method [supporting information (S1) Tables S1 and S2].

The structure of AOS contains two domains with three β-sheets and 15 α-helices (Fig. 2). The major α-domain is predominantly α-helices and contains a conserved structural core around heme, whereas the small β-domain is predominantly β-sheets with two α-helices A' and A in the N terminus, and a 3β helix B.

AOS is structurally similar to other P450s despite the very low sequence identity (Figs. S1 and S2). Superimposing the structures of AOS and human P450s (PDB ID code 2IAG) by using the DaliLite program (34) gave a rmsd of 3.9Å with 401 residues aligned and 14% sequence identity. Superimposing the structures of AOS and bacterial class I P450s (class II: PDB ID code 1OG5) gave a rmsd of 3.6Å with 394 residues aligned and 12% sequence identity. Interestingly, superimposing the structures of AOS and bacterial class I P450BM-3 (PDB ID code 2HPD) gave a lower rmsd (3.0Å) with 408 residues aligned and 13% sequence identity. The differences between AOS and PGIS and other P450s are observed widely (Fig. 3 and Figs. S1 and S2), and the distinct structural features of AOS are mainly related to its unique heme-binding mode, substrate recognition, and specificity (see below).

The native AOS structures in the two different crystal forms were nearly identical with a rmsd of 0.33Å with 466 Cα atoms aligned. The major differences observed include a loop region between β3–2 and β3–3, and the five N-terminal residues that are well defined in the small-cell crystal form (i.e., Native 2), but disordered in the crystal form 1 with a large-unit cell. The AOS molecules are packed more tightly and much better ordered in the small-cell crystal form with a much lower average temperature factor (20.9Å2), whereas the average value was 42.9Å2 for the AOS molecule in the large-cell crystal form.

The structure of AOS bound with substrate analog is also nearly identical to that of native AOS, with a rmsd of 0.22Å over all Cα atoms. This difference is even smaller than that between structures in different crystal forms, indicating that substrate does not induce a significant conformational change in AOS.

Binding Mode with Heme. Unique I-helix. The heme prosthetic group is located mainly between helices I and L in the AOS structure (Figs. 2 and 3). Cysteine-426 acts as the fifth ligand for the iron of the heme cofactor. The classic P450 monooxygenases contain a signature sequence (A/G-G-X-D/E-T-T/S) in helix I, regarded as an oxygen-binding motif with the conserved glycine pointing at the center of the heme and the conserved threonine pointing to the oxygen-binding site. AOS does not require molecular oxygen and contains a unique I-helix different from other P450s (Fig. 3B), with a distinctive sequence (A376-T277-X-G279-G280-X) in its central region. A threonine and glycine (Thr-277 and Gly-280 in AOS) are present at the positions corresponding to the conserved glycine and threonine of bacterial and mammalian P450s, and an asparagine (i.e., Asp-276 in AOS) is present at the corresponding position of the conserved alanine/glycine (e.g., Ala-297 in human P450 1C9; Fig. 3B). In the AOS structure, the residue Gly-280 is ≈4Å from the vinyl group of the heme pyrrole ring A. Thr-277 is <3.5Å from the methyl and vinyl groups of heme pyrrole ring B, and the side chain of Asn-276 is <4Å from heme pyrrole ring C. The threonine, glycine, and asparagine residues are highly conserved in AOSs and the CYP74 family (Fig. S3). The adjacent residues Phe-275 and Lys-282 in the N and C termini are also highly conserved. This sequence (F-N-T/S-X-G-G-X-K) may be regarded as a signature for this family and plays very important roles in heme binding and catalytic function.

Compared with other classes of P450s, the helix I in AOS has an extra amino acid (Cys274 in our alignment, Fig. S1) in the
middle, and the central part of the helix is distorted with a disturbed hydrogen bond pattern. With this extra amino acid, the adjacent residues may be further extended to the active site.

**Unusual heme-binding loop.** The heme-binding loop is a distinct structural feature of AOS related to its unique heme-binding mode. It is located on the proximal face of the heme. In other P450s, this loop usually forms a compact structural feature of AOS related to its unique heme-binding S1), the corresponding loop (S411–G428) in AOS is unusually long middle portion of the heme-binding loop (Fig. 3)

Lys-424 form hydrophobic interactions with the heme and are interacting with the heme iron as the fifth ligand, Ala-427 and corresponding residues in human P450 2C9. Besides Cys-426 residue (lysine) and two identical residues cysteine and glycine with human P450 2C9, PGIS, and other P450s (Fig. 3)

The region between “meander” and the heme-binding loop is also different between AOS and other P450s. Following the conserved meander, AOS contains a 3(10) helix (i.e., G(299)–Y(407)), whereas PGIS has a small β-sheet.

There are other unique interactions between enzyme and heme in the AOS structure. The propionate group of the heme pyrrole ring C also forms charge-charge interactions with His-119 and Lys-123 in helix C. The propionate group of pyrrole ring D interacts with Lys-88. Including the Lys-424 mentioned above, there are four basic residues forming strong charge-charge interactions with heme. These strong and extensive interactions may make the heme group very stable and more restricted in the active site. For other P450s, there are usually only one or two charge-charge interactions formed between enzyme and heme.

All of these structural features define a heme-binding mode for AOS, which is quite different from that of PGIS and other classes of P450s.

**Substrate-Binding Site and Access Channels.** The substrate analog 13(S)-HODE is very similar to the substrate 13(S)-hydroperoxy-9(Z),11(E)-octadecadienoic acid [13(S)-HPOD], with a hydroxyl group in the C13 position, in place of the hydroperoxyl group of the substrate (Fig. 1). In the structure of AOS, the 13(S)-HODE molecule was located in a very narrow and deep pocket observed on the distal side of the heme (Fig. 4 A and B and Fig. S4). This substrate-binding pocket is formed by helices F and I, and loops between helix C and β1-5, between helices F and F’, between helix K and β1-4, and between β3-2 and β3-3, and consists mainly of hydrophobic residues such as Phe-92, Phe-275, and Phe-278. Interestingly, a polar residue (Asn-276) is present in the active site and is very close to heme. This asparagine residue is conserved in all AOSs (Fig. 3) but not in other P450s (Fig. 3B and Fig. S1), suggesting a key role specific for AOS catalysis and function.

The loop F/F’ and the loop between β3-2 and β3-3 are very close to each other on the surface of the substrate-binding pocket, and they define two entrances and channels in both sides, connecting the active site to the surface (Fig. 4 A and B). One channel, designated as channel 1, is mainly between helices F and I. Another channel, channel 2, is between helix F and strand β1-4 with helix A in the entrance. The long fatty acid chain of 13(S)-HODE bends at the C13 position with the 13-hydroxyl group pointing to the heme iron; the long α-chain with the carboxyl group is in channel 1, and the short ω-chain with the methyl group is in channel 2.

Access channels have been observed in other P450 structures in quite different locations and formations. For P450cam, there is one long deep access channel in the open form structure (PDB ID 1K2O) (35), and the channel may be covered in the closed form (e.g., 1DZ4). In zebrafish PGIS, two access channels were observed (32) but in totally different locations; channel 1 is in a near opposite direction with its entrance near helix B’ and the loop B’/C on the other side of the surface, compared with channel 1 in AOS. Channel 2 is also different with an entrance near the helix A’, and β-sheet 1, whereas the entrance of access channel 2 in AOS is near the loop A’/A and the loop between β3-2 and β3-3. Thus, AOS possesses two special access channels quite different from those in other P450s.

**Substrate Recognition.** Six putative substrate recognition sites (SRS1–6) for P450s have been proposed based on the analysis of the CYP2 family and P450cam structure (36). The substrate-binding pocket in AOS may also be classified into similar.
substrate recognition sites although it is quite different from other P450s in sequence and structure.

The loop region before helix C was identified as the SRS1 site in other P450s. In the AOS structure, the corresponding region is a very long loop between strand β1–5 and helix C with 10 more amino acids than in PGIS. The N-terminal portion near Thr-95 is closer to the β-sheet 1, and the regions near Phe-92 and Ser-110 are closer to heme and extend further into the substrate binding pocket, whereas PGIS contains a short helix in the corresponding regions that is further away from the heme. In the AOS–substrate analog structure, residues Phe-92 and Thr-93 are close to the short ω-chain of 13(S)-HODE (Fig. 4A). Ser-110 forms a hydrogen bond with Asn-276, which may interact with the 13-hydroperoxyl group if the true substrate is present in the same position (Figs. 4A and 5), suggesting a critical role in catalysis. The unique I helix in AOS is very important for AOS function, especially its signature sequence region (F275–K282) that is involved in heme binding, substrate access, and catalysis.

The loop after helix K and the N-terminal portion of strand β1–4 are regarded as the SRS5 site in other P450s. This region in AOS contains four prolines and one valine and is close to the short ω-chain of 13(S)-HODE with a distance of ~4 Å from Val-344 and Pro-346. This region is also close to the heme pyrrole rings A and D.

A long loop linking β3–2 and β3–3 is present at the entrance of the active site of AOS. The corresponding region in other P450s is usually even longer, may be present as a small β-sheet 4 that protrudes into the active site and has been regarded as SRS6. In AOS, this loop has a deletion of 14 residues and is nearly located on the molecular surface. Two hydrophobic residues, Pro-458 and Leu-459, point to the long fatty acid chain of 13(S)-HODE, providing a hydrophobic environment. The temperature factor of this loop is very high in all AOS structures, and the loop is nearly disordered in the Native 2 crystal form, suggesting flexibility in this loop region and a possible minor conformational change during substrate binding and product release. This distinctive loop in AOS, together with the loop F/F′, defines the entrances for substrates to access the active site through the channels described above.

The C-terminal region of helix A′ and the N-terminal region of helix A as well as the loop between helices A′ and A are part of the entrance to channel 2. A basic residue (Arg-36) on helix A is present at the entrance. This residue or region may also be involved in substrate recognition and/or product release. All of these SRSs are highly conserved among AOSs, CYP74A, and CYP74C families (Fig. S3), except for rice (Oryza sativa) AOS, which has five fewer residues in the SRS6 site and may
have a more open entrance. These SRSs are quite different in other P450s, including bacterial and mammalian P450s (Fig. S1), and they are highly variable in both sequences and structures, indicating their distinct substrate specificities and functions.

**Implications for Substrate Specificity and Catalytic Mechanism.** In the structure of the AOS complex with substrate analog, the hydroxyl group at the C13 position of 13(S)-HODE is ~7 Å from the heme iron. The substrate 13(S)-HPOD may be situated in a similar position, but should be closer to the heme iron. In a model with the substrate replaced in the active site, the substrate hydroperoxyl group is ~2.4 Å from the heme iron (Fig. 5). Asn-276 in the active site near the heme may interact through a hydrogen bond with the substrate’s hydroperoxyl group on position C13. Ser-110 forms a hydrogen bond with Asn-276. This Asn-276 and the related hydrogen network may play an important role in catalysis and substrate binding.

The basic residue Lys-282 located in the center of helix I and at the entrance of the active site may form a charge–charge interaction with the carboxyl group in the long α-end of the fatty acid substrate and is likely a key residue for substrate binding and recognition. Ser-199 forms a hydrogen bond with Lys-282 and may also interact with the substrate and play a role in substrate binding. To explore the roles of these residues in catalysis and substrate binding, mutagenesis studies of AOS will be necessary.

It has been proposed that the position of the hydroperoxyl group in relation to the methyl end is the key for substrate specificity in AOS (11). In the structure of AOS complexes with substrate analog, the methyl end of 13(S)-HODE is extended into the channel 2 and surrounded by residues in the SRS1 (i.e., F92–T93), SRS2 (V206, T209–L210), SRS4 (Ile-283), SRS5 (P343–P346), and SRS6 (P588–V589) (Fig. 4). It is also close to Tyr-29 and Arg-36 in the A’ and A helices. These interactions are mainly hydrophobic but very broad and extensive, indicating the complexity of substrate specificity. 9-AOS and 9/13-AOS in the CYP74C subfamily may also recognize 9(S)-hydroperoxides as substrates. With 9(S)-hydroperoxide placed in the active site of AOS, the methyl end would be extended closely to the residues at the entrance of channel 2, such as Arg-36. For the 9- and 9/13-AOS, an alanine residue is present in the position corresponding to Arg-36 in AOS and may contact or be close to the methyl end of the 9(S)-hydroperoxide.

Similar to other fatty acid hydroperoxide rearrangements, a homolytic cleavage of the hydroperoxide would be the first step in the conversion of 13(S)-hydroperoxide catalyzed by AOS (37). 13(S)-Hydroperoxide binds to the heme iron via the C13 oxygen atom first, and residues Asn-276 and Lys-282 assist the binding and recognition by interacting with the substrate’s hydroperoxyl group and carboxyl group, respectively (Fig. 5). Heme-assisted homolytic scission of the oxygen–oxygen bond gives an alkoxyl radical and an iron–oxo complex. The alkoxyl radical then undergoes cyclization with the C11–C12 double bond, to yield a C11 radical, which is then oxidized by the iron–oxo complex to form a carbocation. Loss of a proton from the carbocation finally gives the allene oxide.

**Methods**

**Protein Expression and Purification.** The coding region of the AOS cDNA from guayule (P. argentatum) (7) was cloned into the pET28b expression vector with a histidine tag included in the C terminus of the protein. Protein was expressed in *Escherichia coli* BL21(DE3) cells and purified with Ni²⁺-nitrilotriacetic acid agarose and a HisPepSepharose 500 HR gel filtration column according to the procedures reported in ref. 33.

Selenomethionine (SeMet)-substituted AOS was prepared by expressing the recombinant protein in E. coli BL834 (DE3) cells (Novagen) grown in M9 minimal medium supplemented with l-SeMet (Sigma) and purified by following the same protocol used for the native protein.

**Cysteinyl and Data Collection.** Cysteinyl of the native AOS was carried out by using the hanging-drop vapor diffusion method, as described in ref. 33. Briefly, two forms of crystals were obtained from 0.2 M (NH₄)₂HPO₄, 50% 2-methyl-2,4-pentanediol, 0.1 M Tris (pH 8.5). The tetragonal crystal form (Native 1) belongs to the space group P4₂₁ with cell parameters a = b = 126.5, c = 163.8 Å, α = β = γ = 90°. A 2.4 Å native dataset from a tetragonal form of crystal was collected.

Because AOS contains a heavy atom Fe, i.e., the heme iron, a native AOS crystal was used to collect Fe anomalous data by using a wavelength of 1.742 Å with an ADSC Quantum 315 CCD detector at the SBC 19ID beaml ine of the Advanced Photon Source (APS), Argonne National Laboratory (Argonne, IL). A SeMet derivative crystal was used to collect Se anomalous data at the Southeast Regional Collaborative Access Team (SER-CAT) 22ID beaml ine in the APS. Because of crystal decay, only one useful SeMet data set was collected at a wavelength of 0.972 Å.

Crystals of AOS complexed with substrate analog were obtained by cocystalization with 0.1 M 13(S)-HODE (Sigma) under the same crystallization conditions used for native protein crystals. The same crystal form with a similar unit cell was obtained, and a 2.6-Å diffraction dataset was collected with a RuXHIII-imaging plate detector and RuXH x-ray source.

Another native crystal form (Native 2) with a small unit cell (a = b = 113.5 Å, c = 163.8 Å, α = β = γ = 90°) was obtained by dehydration treatment. A high-resolution dataset at 1.8 Å from this crystal form was collected at the SER-CAT 22ID beaml ine in the APS. All datasets were processed by using the program suite HKL2000 (38).

**Structure Determination and Refinement.** The structure of AOS was determined by using the MIRAS method. Besides the Se and Fe anomalous data, collected at wavelengths of 0.972 Å and 1.742 Å, the 2.4-Å data collected from a native crystal by using the RuXHIII x-ray source at a wavelength of 1.5418 Å was used as another Fe anomalous dataset. The program SHARP (39) was used to locate and refine sites of the Se and Fe atoms, yielding an overall figure of merit of 0.22. The phase calculation and density modification were also carried out by using SHARP. Autotracing with the program ARP/wARP (40) was also performed and generated a model containing 377 aa corresponding to 80% of the protein molecule. Further interactive model building and crystallographic refinement were carried out with a native dataset at 2.4 Å by using the programs COOT (41) and CNS (42), respectively.

The structure of the complex of AOS with the substrate analog 13(S)-HODE were determined at 2.6 Å by the difference Fourier method by using the native structure determined above as a template.

The structure in the Native 2 crystal form with small-unit cell was determined at 1.8 Å by the molecular replacement method with the program PHASER (43), with the structure determined above as a search model.

B factors were refined individually for two AOS native structures at 1.8 and 2.4 Å, and grouped B factor refinement was performed for the structure of AOS complexed with 13(S)-HODE at 2.6 Å. Water molecules were added with ARP/wARP (40) and checked manually for inclusion. In the models, the first five amino acid residues in the N terminus were not observed in AOS–analog structure and the native 1 structure. The last five residues in the C terminus were not observed in AOS–analog structure and the native 1 structure. The last five residues in the C terminus were not observed in AOS–analog structure and the native 1 structure. The last five residues in the C terminus were not observed in AOS–analog structure and the native 1 structure.


Fig. S1. Structure-based sequence alignment of allene oxide synthase (AOS) and human prostacyclin synthase (PGIS), human P450 2C9, and bacterial P450BM-3. The secondary structure elements observed in the AOS structure are shown above the alignment. The P450 signature sequence and heme-binding motif are enclosed in green boxes a and b, respectively. Conserved residues are highlighted. This figure was produced with ENDS{[script]\textsuperscript{script}} [Gouet P, Courcelle E (2002) ENDS{[script]\textsuperscript{script}}: A workflow with web interface to display sequence and structure information. Bioinformatics 18:767–768].
Fig. S2. Stereo diagram showing the superimposition of the structures of AOS (cyan) and human PGIS (magenta; PDB ID: 2IAG).
Fig. S3. Sequence alignment of AOSs from guayule (Parthenium argentatum), barley (Hordeum vulgare), rice (Oryza sativa), flaxseed (Linum usitatissimum), tomato (Lycopersicon esculentum), and Arabidopsis thaliana; CYP74C10 from potato (Solanum tuberosum) and CYP74C3 from L. esculentum. The N-terminal transit sequences were omitted for flax AOS, LeAOS, AtAOS, StCYP74C10, and LeCYP74C3. The P450 signature sequence and heme-binding motif are enclosed in green boxes a and b. Conserved residues are highlighted.
Fig. S4. $2F_o - F_r$ electron density omit map of substrate analog 13(S)-HODE contoured at 1.0 $\sigma$. 
### Table S1. Data collection and phasing statistics

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Numbers in parentheses are for the highest resolution shell.

\[ R_{sym} = \frac{\sum_{i} |I_i - \langle I \rangle|}{\sum_{i} |I_i|} \]

where \( I \) is the observed intensity and \( \langle I \rangle \) is the average intensity from observations of symmetry-related reflections. A subset of the data (10%) was excluded from the refinement and used to calculate the free \( R \) value (\( R_{free} \)). \( R \) factor = \( \frac{\sum |F_o - F_c|}{\sum |F_o|} \).
Table S2. Refinement statistics

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Native 1</th>
<th>Native 2</th>
<th>w. 13(S)-HODE</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R$ factor, %</td>
<td>19.8</td>
<td>18.0</td>
<td>21.9</td>
</tr>
<tr>
<td>$R_{free}$ %</td>
<td>24.3</td>
<td>20.8</td>
<td>27.0</td>
</tr>
<tr>
<td>No. of protein atoms</td>
<td>3,742</td>
<td>3,782</td>
<td>3,742</td>
</tr>
<tr>
<td>No. of solvent atoms</td>
<td>206</td>
<td>378</td>
<td>147</td>
</tr>
<tr>
<td>No. of ligand atoms</td>
<td>43</td>
<td>43</td>
<td>64</td>
</tr>
<tr>
<td>Average $B$ factors, Å$^2$</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All atoms</td>
<td>42.9</td>
<td>20.9</td>
<td>27.2</td>
</tr>
<tr>
<td>Protein</td>
<td>43.2</td>
<td>19.4</td>
<td>27.4</td>
</tr>
<tr>
<td>Solvent</td>
<td>40.7</td>
<td>36.0</td>
<td>24.4</td>
</tr>
<tr>
<td>Ligands</td>
<td>34.3</td>
<td>14.8</td>
<td>22.9</td>
</tr>
<tr>
<td>Rmsd from ideal values</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Bond length, Å</td>
<td>0.006</td>
<td>0.005</td>
<td>0.009</td>
</tr>
<tr>
<td>Bond angle, °</td>
<td>1.40</td>
<td>1.30</td>
<td>1.40</td>
</tr>
</tbody>
</table>