Crystal Structure of CYP24A1, a Mitochondrial Cytochrome P450 Involved in Vitamin D Metabolism

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Cytochrome P450 (CYP) 24A1 catalyzes the side-chain oxidation of the hormonal form of vitamin D. Expression of CYP24A1 is up-regulated to attenuate vitamin D signaling associated with calcium homeostasis and cellular growth processes. The development of therapeutics for disorders linked to vitamin D insufficiency would be greatly facilitated by structural knowledge of CYP24A1. Here, we report the crystal structure of rat CYP24A1 at 2.5 Å resolution. The structure exhibits an open cleft leading to the active-site heme prosthetic group on the distal surface that is likely to define the path of substrate access into the active site. The entrance to the cleft is flanked by conserved hydrophobic residues on helices A′ and G′, suggesting a mode of insertion into the inner mitochondrial membrane. A docking model for 1α,25-dihydroxyvitamin D3 binding in the open form of CYP24A1 that clarifies the structural determinants of secosteroid recognition and validates the predictive power of existing homology models of CYP24A1 is proposed. Analysis of CYP24A1’s proximal surface identifies the determinants of adrenodoxin recognition as a constellation of conserved residues from helices K, K″, and L that converge with an adjacent lysine-rich loop for binding the redox protein. Overall, the CYP24A1 structure provides the first template for understanding membrane insertion, substrate binding, and redox partner interaction in mitochondrial P450s.

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Introduction

Mitochondrial P450 enzymes catalyze highly specific reactions in the biosynthesis and degradation of hormones.1,2 Mitochondrial P450s, that is, cytochrome P450 (CYP) 11, 24, and 27 families, cluster into a single evolutionary clan; CYP24A1 itself has less than 40% sequence identity with other P450s.3 An N-terminal leader sequence targeting these P450s to the mitochondrion is cleaved during import; the mature enzymes bind monotopically to the matrix side of the inner membrane. Mitochon-
processes associated with calcium homeostasis, cellular growth, and the maintenance of heart, muscle, immune, and skin function.\textsuperscript{12,13} Evidence suggests that vitamin D insufficiency, due to tissue-specific CYP24A1 overexpression, contributes to development of diabetes\textsuperscript{14} and cancers of the prostate, breast, colon, and lung.\textsuperscript{15–18} A number of promising vitamin D analogs have been developed,\textsuperscript{19} but their administration often induces hormonal resistance. Therefore, a complementary strategy for combating vitamin D insufficiency is to develop specific CYP24A1 inhibitors to retard degradation of the endogenous hormone.

The primary circulating form of vitamin D\textsubscript{3} (25-hydroxyvitamin D\textsubscript{3}) is converted to calcitriol by mitochondrial CYP27B1 (the 25D\textsubscript{3}-1\textalpha-hydroxylase) under hypocalcemic conditions. Calcitriol elicits pleiotropic effects, in part via the vitamin D nuclear receptor (VDR). The CYP24A1 promoter contains multiple vitamin D response elements that respond to VDR signaling; CYP24A1 expression results in side-chain cleavage and elimination of circulating 25-hydroxyvitamin D\textsubscript{3} and 1\textalpha,25-(OH)\textsubscript{2}D\textsubscript{3} by catalyzing hydroxylation at the 23- and 24-positions.\textsuperscript{12,20} The mechanism of CYP24A1’s six-step oxidation pathway for production of calcitriol remains poorly understood.\textsuperscript{21,22} Structural knowledge of CYP24A1 would elucidate the basis for secosteroid specificity and facilitate the rational design of inhibitors, an approach complementary to the synthetic analog design already exploiting structural studies of the VDR.\textsuperscript{23} Here, we report the crystal structure of the rat CYP24A1 determined to 2.5 Å resolution.

**Results**

We developed a bacterial overexpression system for mature rat CYP24A1 lacking the mitochondrial import signal (Δ2–32) to enable biochemical characterization of the purified enzyme.\textsuperscript{24} Subsequent affinity-labeling studies identified a stabilizing mutation for recombinant CYP24A1 (Ser57-Asp) that does not alter catalytic function,\textsuperscript{25} and site-directed mutational analysis using P450 homology modeling established functional roles for conserved residues involved in substrate recognition and catalysis.\textsuperscript{26} Recombinant preparations (Δ2–32, S57D) were solubilized from bacterial membranes and purified in the presence of 3-[3-Cholamidopropyl-dimethylammonio]-1-propane sulfonate (Chaps) (0.8% w/v) by affinity chromatography using Adx linked to Sepharose, yielding monodisperse samples that crystallize readily by vapor diffusion (Supplemental Methods). Detergent exchange from Chaps into CYMAL\textsuperscript{6–5} afforded crystals of CYP24A1, which diffracted to 3.0 Å resolution in space group C\textsuperscript{2} on Stanford Synchrotron Radiation Lightsource (SSRL) beamline 9-2. The structure was solved by molecular replacement with Phaser\textsuperscript{27} using an

**Fig. 1.** The crystal structure of CYP24A1. (a) The structure of CYP24A1 refined at 2.5 Å resolution (R=0.206, R\textsubscript{free}=0.252) shown colored by B-factor from high (red) to low (violet) temperature. P450 structural elements (α-helices, β-sheets) are labeled. Conserved residues from helices F (M245 and F249), G (H271, W275, and F279), and B’ (W134, Y137, and R138) participate in an “aromatic cluster” centered on a water molecule bound to R138 that promotes a membrane-directed (pw2a) substrate access channel. (b) The distal surface of CYP24A1 is shown colored from negative (cyan) to positive (yellow) hydrophobicity (GES scale),\textsuperscript{41} revealing the open channel between the A’–A helix and F–G loop regions. Chaps molecules (pink sticks) from the crystal structure are overlaid to illustrate a model for substrate diffusion from the membrane-associated channel (pw2a) to a putative exit channel (pw3) between the D–E helix region and the E–F loop.\textsuperscript{36}
ensemble of microsomal P450 structures (Supplemental Methods). A model with two molecules in the asymmetric unit was built using Coot28 and MiFit29 and refined using the CCP4 suite.30 Concurrently, an improved “mixed-detergent” method for growing CYP24A1 crystals that eliminated the need for CYMAL®-5 exchange by utilizing a second detergent together with Chaps was developed. CYP24A1 in mixed-detergent micelles with Chaps (0.5% w/v) and common detergents (e.g., CYMAL®, Foscholine®-12) at 0.5- to 2-fold their critical micelle concentration grew readily but diffracted no better than 4 Å resolution.

However, a novel class of detergents, termed facial amphiphiles (FAs),31 are effective in improving CYP24A1 crystal quality in combination with Chaps; single rod-like crystals grown in the presence of Chaps and the FA, 231-chol (3α-hydroxy-7α,12α-bis[(β-D-maltopyranosyl)ethyloxy]cholane), show diffraction to 2.0 Å resolution. A 7.6-fold redundant 2.5-Å data set was collected on SSRl beamline 12-232 and used to complete and refine the CYP24A1 model, including residues 51–514, using the CCP4 suite30 and CNS33 (Table S1). Residues 33–50 of the S57D construct were not visible in the electron density. The two copies of CYP24A1 in the asymmetric unit are very similar with an RMSD between Cα atoms of 0.62 Å.

The open form structure of rat CYP24A1 displays the canonical P450 fold, including the 12 α-helices (A–L) and four β-sheet systems (β1–β4), as well as additional helices A’, B’, and G’ on the distal surface, and K’ and K” between β2 and the conserved heme binding motif (Fig. 1a and Fig. S1). The F-helix of CYP24A1 extends for 18 residues and lacks a well-defined F’-helix seen, for example, in structures of microsomal CYP3A4.34,35 The structural elements defining the substrate-binding cavity, including the β1 and β4 sheets, the B–C loop, and helices E, F, G, I, and K surrounding the heme, exhibit lower-than-
average $B$-factors (Fig. 1a). The B–C loop contains a 3-turn B′ helix that packs closely with helices G and G′. A distinct arrangement of aromatic residues from helices B′ (Trp134 and Tyr137), F (Phe249), and G (His271, Trp275, and Phe279) cluster together with extensive interactions (Fig. 1a, inset), contributing to the lower $B$-values in this region of the structure. The interaction between residues of helix G with helix B′ entraps a water molecule hydrogen bonded with the carbonyl oxygen of Trp134 and the side chain of Arg138; the latter residue engages in a

The Crystal Structure of CYP24A1

The substrate access channel apparent in CYP24A1 is occupied by detergent molecules (Fig. 1b and Fig. S4) and lies among the B–B′ segment of the B–C loop, the β1 sheet, and the F–G loop (pw2a); this channel appears to partially merge with a secondary channel that also lies between the B–B′ loop and the β1 sheet and enters from below the F–G loop (pw2b). A similar substrate access channel, oriented toward the membrane, is present in the structure of the microsomal P450, CYP46A1, which is a cholesterol 24-hydroxylase. This feature distinguishes CYP24A1 from the nonspecific microsomal vitamin D 25-hydroxylase, CYP2R1, which contains a pw2c substrate-binding channel typical of enzymes from the CYP2 family.

Among currently available structures, the overall topology of structural features that form the substrate-binding cavity in CYP24A1 is most similar to that of CYP3A4. A comparison of the two structures suggests that the open cleft observed for CYP24A1 results from the close packing of helix F with helix G and the elevated pitch of both helices that separates helix F from the β4 sheet. This separation is increased by the conformation of the β1–4 sheet region, which is similar to that seen in CYP3A4. This conformation exposes a larger portion of the heme surface and a widening of the substrate-binding cavity close to the catalytic center, relative to structures of mammalian P450s of family 1 or family 2. In structures of CYP3A4, helix F does not extend over the substrate-binding cavity and the polypeptide chain exhibits a coil structure as it passes across the cavity and connects with helix F′. In contrast, the helix F of CYP24A1 extends across the structure toward helix G′ without a well-defined helix F′.

Analysis of the CYP24A1 structure identified hydrophobic surfaces of helices A′ and G′ as potential anchors to the mitochondrial inner membrane; together, these short helices flank the substrate access channel (Fig. 1a and b). The corresponding A′–A-helix and F–G loop regions of microsomal P450s are also implicated in membrane association. Computational prediction of CYP24A1 insertion into a simulated lipid bilayer confirms this analysis (Fig. 2a and b). The predicted protein:membrane association transfer free energy (−9.0 kcal/mol) is within the range of values predicted from other P450 crystal structures [e.g., CYP3A4 (1TQN) = −20.7 kcal/mol and CYP46A1 (2Q9F) = −4.7 kcal/mol]. This analysis also suggests that residues on helices A′ and G′ can penetrate approximately 7 Å into the alkyl chain region of the bilayer, or up to 22 Å into the outer leaflet (Fig. 2a).

Alignment of the A′ and G′ segments with other mitochondrial P450s identifies each as a membrane insertion sequence (MIS), implying a conserved membrane binding architecture within mitochondrial CYPs 11, 24, and 27 (Fig. 2c and d). In MIS-1, the conserved proline-rich (PGP) region is followed by a hydrophobic segment of 9–12 residues bracketed by aromatic amino acids (Trp64 and Trp75, Fig. 2c). MIS-2 encompasses the amphipathic G′-helix, a distinctive feature of the structure (Fig. 1a), in which polar residues are oriented back toward the protein and hydrophobic residues point outward. MIS-2 is delimited by a conserved proline in the F–G loop region (Pro256) and a tryptophan at the N-terminus of the G-helix, present in all mitochondrial P450s (Trp268, Fig. 2d). The alignment in Fig. 2d suggests that the amphipathic nature of the G′-helix is a conserved feature of mitochondrial P450s. A number of conserved, basic residues are also arrayed in the vicinity of the A′ and G′ hydrophobic regions, consistent with interactions with phospholipid head groups (Fig. S3).

Four molecules of Chaps were observed in copy A of the 2.5-Å structure of CYP24A1. Two molecules are associated with membrane binding regions (MIS-1 and MIS-2) and structural lattice contacts; one is bound in the substrate-access channel and a fourth is positioned above the heme in a nonproductive binding orientation (Fig. 1b). Alternatively, two molecules of CYMAL®-5 occupy the pw2a access channel in the independent 3.0-Å structure of CYP24A1 (Fig. S4). The arrangement of either Chaps or CYMAL®-5 in isomorphous CYP24A1 structures suggest that the open form of CYP24A1 is a biologically relevant conformation that is not dependent on the specific details of detergent binding. Additionally, the presence of trapped detergent molecules is consistent with the hydrophobic nature of the access channel and binding pocket and illustrates a potential pathway for lipophilic substrates to transit from the membrane to the active site.

There is well-defined electron density for a Chaps molecule (CPS-600, 3K9V) close to the heme within the substrate-binding pocket of CYP24A1 (Fig. 3), but density for the zwitterionic side chain extending out from the cavity is weaker. The hydrophobic face of the cholate ring is oriented toward the heme surface and nonpolar residues of helices B′ and F, whereas the polar face of Chaps is oriented toward
polar protein groups, allowing three hydrogen bonds to be formed. In solution, Chaps does not inhibit the catalytic function of CYP24A1 and does not induce a Type I spectral perturbation of the heme typical of a proper substrate.24–26 Hence, while detergent association in the active site appears strong, it likely represents a nonproductive interaction that does not promote the enzyme’s closed form. Nevertheless, Chaps binding is illustrative of the hydrophobic nature of the active site and its ability to provide specific interactions with amphipathic molecules.

Residues surrounding the active-site cavity are contributed from nine regions of the CYP24A1 fold: B–B′ region (Leu129 and Ile131), B′-helix (Trp134), B′–C region (Met148), F-helix (Met245, Met246, and Phe249), G-helix (His271 and Trp275), I-helix (Leu325, Ala326, Glu329, and Thr330), K-helix/β1–4 loop (Val391), β1–4 sheet (Phe393, Thr394, and Thr395), and the β4-1/β4-2 turn (Gly499 and Ile500) (Fig. 3). Mutational analysis of CYP24A1 based on P450 homology modeling studies has attributed roles in substrate binding and catalysis for 13 of the residues identified in the structure: Ile131, Trp134, Met148, Met245, Met246, Phe249, Ala326, Glu329, Thr330, Val391, Thr394, Gly499, and Ile500.26,48,49 Structural analysis also supports predicted roles for Ala326 (I-helix) and Ile500 (β4-1/β4-2 loop) in modulating regiospecificity of hydroxylation in CYP24A1.50,51 A residue from the β1–3 strand (Thr416) is also linked to this process50 but is found outside the Chaps binding site in the open channel. However, this location could be relevant for secosteroid binding by analogy to cholesterol-3-sulfate binding in CYP46A1.37

A hybrid homology model for CYP24A1 based on the structures of mammalian CYP2C5 and bacterial CYP102 successfully predicted multiple residues in the CYP24A1 binding pocket, including Met246, Phe249, Val391, Thr394, and Ile500.26 However, this
model does retain some template bias as it predicts substrate access and binding similar to CYP2C5, via the solvent-accessible pw2c channel that is occluded in the structure. A P450 structural-motif-based method of homology model building utilized by Masuda et al. appears to be highly effective at predicting key elements of the CYP24A1 tertiary structure.48 This 3D model has been used to make strong predictions concerning the positioning, and putative role, of conserved residues in the CYP24A1 active site, including Ile131, Trp134, Met148, Met246, Ala326, and Gly499. Flexible ligand docking simulations with Chaps (as a control) and 1α,25-(OH)2D3 were conducted against the rat CYP24A1 structure (Table S2) using Autodock 4.0 (see Materials and Methods) to clarify the functional role of putative substrate-binding residues in the open form.70–72 As illustrated in Fig. 4, 1α,25-(OH)2D3 docks in the open form of CYP24A1 with computed nanomolar affinity (Ki = 2.65 nM) in a reproducible conformation stabilized by two hydrogen bonds [between the 3-OH group of the vitamin D A-ring and the B–B′ loop (Leu129), and the 25-OH of the side chain and the I-helix (Leu325)] and multiple hydrophobic interactions among key conserved residues (Ile131, Trp134, Met246, Phe249, Ala326 Val391, Thr394, Thr395, Gly499, and Ile500). The docking calculations confirm that the open form of CYP24A1 is well organized to bind 1α,25-(OH)2D3, in accord with predictions made by homology modeling concerning residues likely to bind the A–C–D ring system (Leu129, Ile131, Trp134, Phe249, Thr394, and Val391) or side chain (Leu325, Ala326, Met246, and Ile500) of the vitamin D hormone. In this configuration, the secosteroid is well positioned to interact with the aromatic cluster of residues among helices B′, F, and G via interactions with the B–C loop (Leu129, Ile131, Met246, and Met448) and helix F (Met246 and Phe249). However, it is unlikely that this result represents the

![Fig. 4. Secosteroid docking in the open form of rat CYP24A1. A model for 1α,25-(OH)2D3 (yellow) binding in the heme-centered (pink) active site of CYP24A1 was developed with a crystal structure-calibrated docking protocol, using Autodock 4.0 (see Materials and Methods). Amino acid residues that flank the secosteroid docking site are shown from the B–B′ loop (L129 and I131), the B′–C loop (M148), helix F (M246 and F249), helix I (L325, A326, E329, and T330), the K-helix/β1-4 loop (V391 and F393), the β1–4 sheet (T394 and T395), the β1–3 sheet (T416), and the β4-1/β4-2 turn (G499 and I500), and notable hydrogen bond distances (in angstroms, red) are given with respect to the computed H atom positions. Individual carbon atoms on 1α,25-(OH)2D3 are labeled for reference. Multiple hydrophobic interactions and two hydrogen bonds between the 3-OH group and the B–B′ loop (L129) and the 25-OH group and helix I (L325) are predicted to stabilize secosteroid binding in the open form. In this configuration, the C21-methyl group superimposes with a structural water (WAT10) bound to the heme (shown as red-dotted sphere); this water molecule was excluded from the calibrated docking experiment shown here (Table S2; Fig. S5). An alternative docking model derived using control parameters, which include WAT10, is presented in Fig. S6.](image-url)
substrate’s terminal binding configuration relevant for the catalytically active enzyme; the C21-methyl group of 1α,25-(OH)2D3, and not the target C23 or C24 carbons, is positioned over the heme iron, where it is superimposed with an ordered water molecule (WAT10, 3K9V). This result implies that, in the open form, secosteroid substrates bind over the heme in a fashion that brings the C21-methyl group into position to perturb the water-bound heme iron (Fig. 4 and Fig. 5S). Further, when WAT10 is included in the docking simulations, a similar solution for 1α,25-(OH)2D3 is obtained (Table S2; Fig. S6), but here, the C21-methyl is forced to rotate away from WAT10, forming a hydrophobic interaction with the important catalytic residue Ala326; the hydrogen-bonding network is also altered as the 3-OH group is repositioned to interact with the β1–4 loop (Leu129) (Fig. S6). These derived results suggest that electrostatic interactions between the 25-OH group and the l-helix (Leu325), and hydrophobic interactions among the hormone’s C–D ring system and the B–B′ loop (Met148) and the K-helix/β1–4 loop (Val391), are essential for high-affinity substrate binding in the open form. This analysis validates published CYP24A1 homology modeling studies concerning the composition of residues in the active site and supports our hypothesis that the open form is a biologically relevant conformation, useful for studying substrate binding in mitochondrial P450s.

The CYP24A1 structure also reveals structural elements and key residues for interacting with the [2Fe–2S] electron transfer protein, Adx (Fig. 5a). Four basic residues from helices K (Lys378 and Lys382) and L (Arg465 and Arg466), involved in Adx recognition and electron transfer in CYP27B1,52 are juxtaposed on the proximal surface. These basic residues are conserved in mitochondrial P450s but divergent in microsomal forms (Fig. S7). Structural elements on the proximal surface include segments of helices B, C, D, and K, helices K′ and K″ flanking the meander region, and the Cys loop (Fig. 5b), all of which are implicated in redox partner binding52–55 and circumscribe the four basic residues. Compared to microsomal P450s, all mitochondrial CYPs contain a tryptophan (Trp440 in CYP24A1) in the conserved motif PxRWL in the K″-helix (Fig. S7). Trp440 interacts with the K-helix via another conserved residue, Glu383, within the invariant KExxR motif (Fig. 5b and Fig. S7). This conserved tertiary interaction, which anchors the lysine-rich loop (residues 442–446, QKEKK), and is next to Lys378 and Lys382, appears to be important for Adx interaction. For example, it links a substrate recognition sequence (SR55, K-helix/β1–4 sheet)56 with the lysine-rich loop, which mediates redox protein interaction in P450cam.67 It also displays the conserved residue Leu441 on the protein surface, providing a complement to electrostatic interactions in CYP24A1-Adx recognition.

**Discussion**

Mitochondrial CYP24A1 is a monotopic membrane protein that binds tightly to, but does not span, the lipid bilayer.47,58,59 The CYP24A1 structure indicates that CYP24A1 associates with the membrane using hydrophobic MIsS and conserved basic residues surrounding helices A′ and G′ (Fig. 2 and Fig. 5S). Prior studies of both microsomal (CYP2C2 and CYP3A4) and mitochondrial (CYP11A1 and CYP27A1) P450s have established roles for the N-terminal proline-rich motif, A′–A-helix region, and the F–G loop in membrane binding.42–45,60 The
CYP24A1 structure is consistent with these results and clarifies the key role of the conserved membrane insertion elements (MIS) encompassing helices A′ and G′. However, the MIS-1 and MIS-2 features of CYP24A1 are not shared by bacterial P450-BM3 (CYP102), a fatty-acid-metabolizing, soluble P450 that otherwise shares common features with mitochondrial P450s. MIS elements could also mediate protein:lipid interactions, as postulated for other monomeric membrane proteins, such as the C2 domain of cytosolic phospholipase A2. Penetration of CYP24A1 up to 22 Å into the bilayer could account for the roles of specific lipids, for example, cardiolipin, in altering mitochondrial P450 function.

The open conformation of CYP24A1, compatible with substrate diffusion from within the membrane, is uniquely stabilized by a cluster of conserved aromatic amino acids from helices B′, F, and G. The aromatic cluster blocks the solvent-accessible channel (pw2c) formed by helices B′, G, and I and promotes formation of a pw2a channel directed toward the membrane. Stabilization of an open conformation may also contribute to substrate specificity in CYP24A1. Active-site collapse to a closed state, as required to exclude solvent during catalysis, may be triggered by solvent displacement and stabilized by the enthalpy of specific interactions with the biological substrate. Hence, the open form would disallow efficient, nonspecific oxidation of improper substrates that are not able to stabilize the closed state, defining at least one mechanism by which CYP24A1, and perhaps mitochondrial P450s, may achieve their exquisite specificity. Docking analysis supports this hypothesis, as 1α,25-(OH)2D3 is computed to bind deep within the active site where it can interact with the heme group and presumably accommodate the cavity collapse required for catalysis. In contrast, the non-substrate Chaps, which also dock the enzyme with computed nanomolar affinity (K<sub>i</sub> = 5.79 nM), is bound in a shallow conformation with limited access to the heme that would not be predicted to inhibit secosteroid accessibility. These studies provide new insight into the enzyme’s active-site organization before substrate binding, providing testable hypotheses concerning the determinants of secosteroid recognition. These contributions should prove useful in the evaluation and design of vitamin D analogs and CYP24A1-specific inhibitors.

The proximal surface of CYP24A1 interacts with Adx and is rich in basic residues, including the invariant arginine pair (Arg465 and Arg466) at the N-terminal end of the L-helix (Fig. 5a). These side chains protrude below the heme and are within 8–10 Å of the conserved K-helix residues, Lys378 and Lys382. Mutational analysis of the arginine pair in CYP11A1 and CYP27B1 suggested that the first arginine contributes to protein folding events and/or heme binding, as mutations abolish enzyme function and promote the P-420 form. An Arg465 mutant of CYP24A1 (R465F) showed a similar dysfunction. Hence, Arg465 may play a dual role. The second arginine (Arg466) has been more clearly linked to catalytic function and is thought to mediate both electron and oxygen transfer steps. In CYP24A1, Arg466 is positioned to promote electron transfer between Adx and the heme iron via main-chain interactions with Cys462, as modeled for the CYP101:putidaredoxin complex. Conserved residues from the K-helix (Lys378 and Lys382) and the lysine-rich loop are also positioned to interact with the very acidic surface of Adx that surrounds the [2Fe–2S] cluster. The link between the K and K′ helices provided by the conserved Glu383-Trp440 tertiary structure interaction suggests that presentation of the PxRWL motif is another key aspect of Adx recognition.

In summary, we have solved the first structure of a mitochondrial P450, CYP24A1, which defines key features for this important class of enzymes. A membrane-directed (pw2a) substrate access channel is stabilized in an open conformation by an aromatic cluster formed among conserved residues on helices B′, F, and G. Two conserved MISs overlap hydrophobic features on the distal surface near helices A′ and G′. Membrane insertion and orientation of the enzyme’s hydrophobic channel show how a lipophilic substrate, in this case vitamin D, can be transferred from the bilayer to the active site. Conserved surface features from helices K, K′, and L, and the lysine-rich proximal loop of the meander region, define components of the basic Adx binding site. The composition of residues in the active site is consistent with prior biochemical, mutagenesis, and homology modeling data for CYP24A1, and related mitochondrial P450s, and docking simulations with 1α,25-(OH)2D3 clarify the structural determinants of secosteroid recognition in the enzyme’s open conformation.

**Materials and Methods**

Recombinant rat CYP24A1 enzyme (Δ2–32, S57D mutant) was purified using established protocols. Some modifications to our original procedure were required for crystallization (as detailed in Supplemental Methods). In short, a secondary chromatography step was added, which allows CYP24A1 samples prepared in Chaps (Anatrace) to be exchanged into alternative detergents, like CYMAL<sub>6</sub>–5 (Anatrace), as described by Wester et al. Detergent-exchange experiments identified conditions for growing CYMAL<sub>6</sub>–5-based crystals that diffract to 2.8 Å and allowed the CYP24A1 structure to be solved by the molecular replacement method using Phaser. Later, a mixed-detergent method was established for improving CYP24A1 crystal quality without the need for detergent exchange. FA detergents developed at Scripps were found to produce high-order CYP24A1 crystals that diffracted to 2.0 Å resolution when mixed with Chaps. Diffraction data for all CYP24A1 crystal forms were collected at the SSRL (SSRL BL-2 and BL12-2).

Detailed methods for structural determination and refinement, computational (surface and membrane binding) analysis, and flexible ligand docking studies with Autodock 4.0 are provided in Supplementary Materials. All sequence alignments were developed using the program ClustalX2 labeling and format are explained.
in Supplemental Methods. All figures for this manuscript were generated using PyMOL.\textsuperscript{74}

**Accession codes**

The atomic coordinates and structure factors have been deposited in the Protein Data Bank\textsuperscript{†} (PDB codes 3K9V and 3K9Y).

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In Supplemental Methods. All figures for this manuscript can be found, in the online version, at doi:10.1016/j.jmb.2009.11.057\textsuperscript{**}


**Supplementary Data**

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2009.11.057

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